

The Conformational Properties of Methyl α -(2,8)-Di/Trisialosides and Their *N*-Acyl Analogues: Implications for Anti-*Neisseria meningitidis* B Vaccine Design[†]

Austin B. Yongye,[‡] Jorge Gonzalez-Outeiriño,[‡] John Glushka,[‡] Verena Schultheis,[§] and Robert J. Woods^{*,‡}

Complex Carbohydrate Research Center, 315 Riverbend Road, Athens, Georgia 30602, and Theoretische Biophysik, Lehrstuhl für Biomolekulare Optik, Ludwig-Maximilians-Universität, Oettingenstrasse 67, 80538 Munich, Germany

Received March 13, 2008; Revised Manuscript Received August 19, 2008

ABSTRACT: The conformational properties of di- and trisaccharide fragments of the polysialic acid *O*-antigen capsular polysaccharide (CPS) of *Neisseria meningitidis* B (*NmB*) have been investigated by a combination of solution phase NMR spectroscopy and explicit-solvent molecular dynamics (MD) simulations. Simulations employing 100 ns of conventional MD, as well as 160 ns of replica exchange MD (REMD), with the GLYCAM06 force field were shown to be in agreement with experimental NMR scalar *J*-coupling and NOE values. The presence of conformational families has been determined by monitoring interglycosidic torsion angles, by comparing structural superimpositions, as well as via a Bayesian statistical analysis of the torsional data. Attempts to augment the immunogenicity of *NmB* CPS often involve chemical modifications of the *N*-acetyl moiety. Here the effects of these chemical group modifications on the conformational properties of the trisialoside have been probed via REMD simulations of the *N*-glycolyl, *N*-propionyl, *N*-propyl and *N*-butanoyl analogues. Although there were conformational families unique to each non-native analogue, the chemical modifications resulted in largely equivalent overall conformational phase-spaces compared to the native trisialoside. On the basis of the conformational distributions, these shared conformational properties suggest that a recurrent global conformational epitope may be present in both the native and chemically modified CPS fragments. Explanations are therefore provided for monoclonal antibody cross-reactivity, in terms of recognition of a shared global CPS conformation, as well as for lack of cross-reactivity, in terms of fine structural differences associated with the *N*-acyl groups, which may be dominant in highly matured antibody responses.

Neisseria meningitidis (*Nm*¹) is a Gram negative bacterium that is responsible for septicemia and meningococcal disease in neonates (1). Annually, about 500,000 cases of meningococcal meningitis are reported worldwide, with 2600 occurrences in the United States. Ten percent of infections result in deaths, while survivors may suffer from seizures, hearing impairments and brain damage (2). Thirteen strains have been classified serologically, based on the identity of the polymeric repeat units and the types of glycosidic linkages in the *O*-antigen capsular polysaccharides (CPS) that envelop the surface of the bacteria (3). Of these thirteen strains, five strains, A, B, C, W-135 and Y, have been identified as the main causative agents of virulent invasive meningococcal disease, Table 1 (4–7).

[†] The authors thank the National Institutes of Health (Grants RR05357 and GM55230) for funding. Computational time was provided in part by a Grant (#9607) from the Environmental Molecular Sciences Laboratory (EMSL) at Pacific Northwest National Laboratory (PNNL). V.S. thanks the Boehringer Ingelheim Fonds for a Ph.D scholarship.

* To whom correspondence should be addressed. E-mail: rwoods@ccrc.uga.edu. Phone: (706) 542-4454. Fax: (706) 542-4412.

[‡] Complex Carbohydrate Research Center.

[§] Ludwig-Maximilians-Universität.

¹ Abbreviations: *Nm*, *Neisseria meningitidis*; *NmB*, *Neisseria meningitidis* serogroup B; *NmC*, *Neisseria meningitidis* serogroup C; mAb, monoclonal antibody; ISPA, isolated spin-pair approximation; PSA, polysialic acid; Neu5Ac, α -5-*N*-acetylneuraminic acid; CPS, capsular polysaccharide; MD, molecular dynamics; REMD, replica exchange molecular dynamics; *E. coli*, *Escherichia coli*; *N*-Glc, *N*-glycolyl; *N*-Pr, *N*-propionyl; *N*-Prop, *N*-propyl; *N*-But, *N*-butanoyl.

Table 1: CPS Structures of the Five Most Virulent *N. meningitidis* Serotypes

serotype	oligosaccharide repeat units	acetylation
W-135	-6)- α -D-Gal-(1–4)- α -Neu5Ac-(2-	7 and/or 9 <i>O</i> partial (5)
Y	-6)- α -D-Glc-(1–4)- α -Neu5Ac-(2-	9 <i>O</i> partial (5)
C	-9)- α -Neu5Ac-(2-	7 and/or 8 <i>O</i> (6)
B	-8)- α -Neu5Ac-(2-	— (6)
A	-6)- α -D-ManNAc-(1-OPO ₃ -	3 <i>O</i> (7)

The lack of cross-reactivity between anti-CPS sera indicates that, despite the fact that several *Nm* capsules bear striking carbohydrate sequence similarities, they must display unique antigenic determinants or epitopes. That is, each CPS either must have a unique overall 3D conformation or must display its immunodominant regions in unique 3D configurations. For example, serogroups B (*NmB*) and C (*NmC*), which are both linear homopolymers of α -*N*-acetylneuraminic acid (Neu5Ac) containing α -(2,8) and α -(2,9) glycosidic linkages, respectively, elicit antibodies that do not cross-react with each other's CPS (3). It should be noted that variations in the level and position of *O*-acetylation also exist between the serotypes (Table 1) (5–7). It is remarkable that, despite the apparent sensitivity to carbohydrate sequence, cross-reactivity of a human monoclonal antibody (IgM^{NOV}) with the *NmB* CPS and poly(adenine) and other polynucleotides has been observed (8), and attributed to similar conformational properties and spatial distribution of anionic charges between the carboxyl groups of the B polymer and the

phosphate groups of the polynucleotide (9). This result serves to illustrate a recurrent phenomenon, namely, that IgM class antibodies often show broader cross-reactivity than seen with more highly affinity matured IgG antibodies, and it is necessary to consider this factor when discussing CPS reactivities. Each *NmB* CPS may also display differences in the apparent size of the epitope. For example, in binding inhibition assays a minimum of approximately ten residues of the *NmB* CPS was required for effective inhibition of antisera binding to the intact polysaccharide antigen, while for strain C, maximal inhibition occurred with a more conventional hapten length of four or five residues (10). In addition to antigenic variations arising from differences in glycosidic linkage types, epitope size may also be altered by chemical modifications. In the case of *NmB*, inhibition studies indicated that reduction of the carboxylic acids to primary alcohols (sialitols) led to a decrease in the epitope size to approximately six residues (11). The requirement in *NmB* for an oligosaccharide length of at least ten residues, to inhibit antibody binding to the native CPS, has led to the postulation that this CPS adopts an extended conformational epitope (10). Extended epitopes are generally assumed to reflect the presence of a conformational requirement for binding, giving rise to the concept of a conformational epitope. All extended epitopes should therefore be conformational, but given the fact that oligosaccharides often exist in well-defined conformations, even a short sequence may have a conformational requirement for binding. This is in contrast to the case of short peptide epitopes.

Currently, two CPS-based conjugate vaccines against *Nm* are available (12): a bivalent form, composed of groups A and C, and a tetravalent form containing A, C, Y and W-135. A conjugate vaccine against *NmB* is not currently available due to the poor immunogenicity of its CPS, which is structurally equivalent to the polysialic acid (PSA) autoantigen present in the glycoproteins of human neural cell adhesion molecules, and is therefore not immunostimulatory (13). The corollary to this structural similarity is that antibodies that bind to the *NmB* CPS can also cross-react with PSA (14), raising a question as to whether or not a CPS-based vaccine would have undesirable side effects. However, studies of human maternal and placental cord sera have found that "natural" anti-PSA antibodies were present in almost all matched pairs of sera (15); suggesting that increased levels of anti-*NmB* CPS antibodies elicited by vaccination would not be pathologic (15). It has also been hypothesized that the weak immunogenicity of *NmB* CPS-based vaccines arises from the formation of transglycosidic lactone rings, between the hydroxyl group at C9 and the carboxylic acid in the preceding residue (16). Lactonization could alter the antigenicity and immunogenicity of the CPS by changing its conformation and charge distribution (16), and indeed it has been observed that as little as 9% lactone content was sufficient to attenuate the antigenic properties of the CPS (16). Recent NMR studies of the *NmB* CPS displayed on *E. coli* K12 cell surfaces, under physiological conditions, showed no evidence of lactone formation, confirming that it is not a biologically common form of the CPS (17).

Several attempts to develop immunogenic *NmB* vaccines have focused on modifications of the amido group, for example by replacement of the *N*-acetyl with *N*-glycolyl (*N*-

Gc), *N*-propionyl (*N*-Pr), *N*-propyl (*N*-Prop) or *N*-butanoyl (*N*-But) moieties (11, 18–22). Somewhat perplexing cross-reactivities, as well as varying levels of protection and bactericidal activity, have emerged from immunizations with these conjugate vaccines. Several of the modified polysaccharides appear to present an extended epitope similar to that found in *NmB*, as judged by cross-reactivity with monoclonal antibodies known to display specificity for the *NmB* extended epitope. For example, the cross-reactivity of the *N*-Pr CPS with the well-characterized bactericidal *NmB* monoclonal antibody 735 (IgG) indicates that both the *N*-Pr analogue and the native CPS share a similar extended epitope of approximately ten residues (22). However, some CPS analogues, such as the *N*-Prop derivative, bind poorly to mAb 735 (18), suggesting that the *N*-Prop analogue differs significantly from the native CPS, either in overall conformation or in local contacts involving the *N*-acyl groups. Notably, the presence of an extended epitope is not sufficient to ensure cross-reactivity among the modified immunogens and/or antisera. For example, although the *N*-Pr analogue and the native CPS both bind to mAb 735 (22), immunization with the *N*-Pr conjugate, while giving a strong antibody titer against the *N*-Pr CPS, gave little cross-reaction with *NmB* CPS (21). Similarly, a bactericidal monoclonal antibody, IgG 13D9, generated by immunization with the *N*-Pr conjugate vaccine showed no affinity for the native *NmB* CPS (22), although it did cross-react with the *N*-But CPS (23). Subsequent analysis confirmed that mAb 13D9 does bind to the bacterial surface of *Nm*, leading to its bactericidal activity, but does not bind to *NmB* CPS (22). Thus, the bactericidal activity of mAb 13D9 can only be understood by accepting that it recognizes an epitope on the bacterial surface, other than that for which the vaccine was designed (22).

As an alternative to modification of the *NmB* CPS, attempts have been made to stimulate antibody production against both the *NmB* and *NmC* CPS by immunizing with vaccines made from the CPS of *E. coli* K92. The K92 CPS consists of alternating α -(2,8) and α -(2,9) linkages, and conjugate vaccines using this CPS could therefore be expected to produce by default anti-*NmB* (2,8) or *NmC* (2,9) responses (15, 24). Although vaccination with K92 CPS conjugates did yield anti-*NmC* responses, they were poorer than direct immunization with *NmC* conjugate (15). Antisera from K92 vaccinations showed only a very modest if any affinity for *NmB* CPS (15, 21). In an attempt to enhance the immune response to K92 conjugate, following on the results with chemically modified *NmB* CPS, the *N*-Pr analogue of K92 was generated. However, immunization with *N*-Pr K92 conjugate did not lead to any improvement in cross-reactivity, but rather the antisera lost all affinity for both *NmB* and *NmC* CPS (21). In contrast to the case of *NmB*, the *NmC* CPS does not display an extended epitope (21), and the lack of affinity for *NmB* displayed by the *E. coli* K92 antisera has been ascribed to the absence of an extended epitope in K92 (21). The origin of the deleterious effect of *N*-propionylation on the ability of K92 conjugate to elicit anti-*NmC* antibodies remains unclear, particularly since the *N*-Pr K92 antisera retained the ability to recognize native K92 CPS.

The potential relevance of an extended conformational epitope to the antigenicity, immunogenicity and bactericidal activity of *NmB* CPS (10) has stimulated numerous investigations of the conformational properties of this CPS

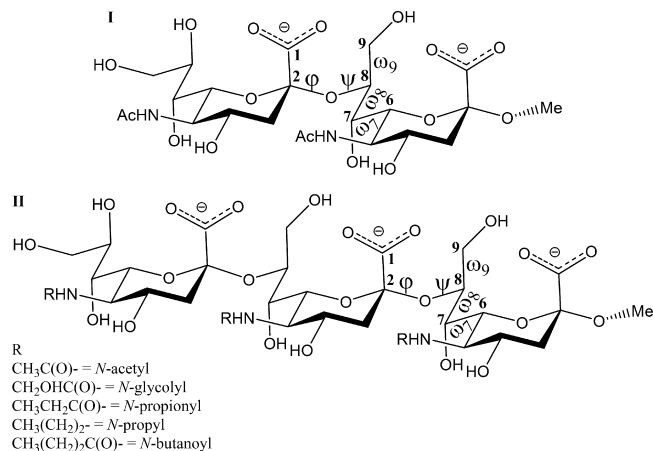


FIGURE 1: Schematic representation of the α -2,8-linked *N*-acetylneuraminic acid dimer (I) and trimer (II). The dihedral angles are defined as follows: $\varphi = \text{C1}-\text{C2}-\text{O8}'-\text{C8}'$, $\psi = \text{C2}-\text{O8}'-\text{C8}'-\text{C7}'$, $\omega_9 = \text{O9}-\text{C9}-\text{C8}-\text{O8}$, $\omega_8 = \text{O8}-\text{C8}-\text{C7}-\text{O7}$ and $\omega_7 = \text{O7}-\text{C7}-\text{C6}-\text{O6}$.

(9, 11, 18, 25, 26). Understanding the structural features that determine the conformational properties that mediate the specificity and antigenicity of interactions between this CPS and bactericidal antibodies could provide a basis for the interpretation of the antigenic properties of related structures and possibly for the rational development of more immunogenic and bactericidal vaccines against *NmB*. Any chemical modification that alters the conformational epitope of a CPS-based immunogen may lead to attenuated antigenicity or unexpected cross-reactivity. By conserving the epitope conformation responsible for generating bactericidal antibodies (the bactericidal epitope) it should be possible to categorize putative immunogens not only in terms of their antigenicity but also in terms of their potential to elicit bactericidal antibodies. However, the flexibility of *NmB* in particular, as well as its homopolymeric sequence, currently prevents a direct resolution of the NMR data into definitive conformational families. The linkage regions between the residues of the *NmB* CPS may be characterized by four conformationally significant rotatable bonds, Figure 1. The presence of these flexible dihedral angles (φ , ψ , ω_7 , ω_8) renders the characterization of the 3-D structure and dynamics of the *NmB* CPS particularly challenging. When conformational variations are significant, such as in carbohydrates, the risk of generating virtual 3D conformations from the use of NMR restraints to guide the structural refinement is significant (27). MD simulations can be employed to provide a structural basis for interpreting the NMR data or for *a priori* investigations of the effects of chemical modifications on the size and shape of the CPS epitopes. For example, MD and NMR studies have been used to elucidate the role of sialic acid residues in determining the overall conformation of the CPS from type III group B *Streptococcus* (28). For the specific case of the *NmB*, molecular mechanics potential energy maps and MD simulations have been reported for the α -(2,8)-disialoside (9, 21) and α -(2,8)-tetrasialoside (26), respectively, in the gas phase. Explicit-solvent MD simulations employing simulation times of 3 ns (21) and 1 ns initiated from nine different low-energy conformations (29) of the α -(2,8)-disialoside and 530 ps for the α -(2,8)-tetrasialoside (26) have also been reported. However, it is now well established that the lifetimes of rotamers in

glycosidic linkage regions can span more than the 10 ns time frame (28, 30). In order to identify all of the significant conformational states, as well as to determine their populations, simulations on the order of 100 ns are typically required (28, 30, 31).

Here, we characterize the conformational properties of methyl α -(2,8)-di- (I) and trisialoside (II) haptens from the CPS of the *NmB*, employing data from 100 ns MD simulations performed with a force field (GLYCAM06) (32) that has been specifically developed for carbohydrate simulations. The theoretical conformations were then independently assessed for their experimental accuracy through comparison with new NMR NOE and *J*-coupling data for I and II. Additionally, replica exchange MD (REMD) simulations were performed for I and II and the *N*-Gc, *N*-Pr, *N*-Prop and *N*-But analogues of II, Figure 1. The REMD protocol was recently shown to lead to excellent agreement with experimental populations for simulations of glycerol (33), suggesting that this approach may be appropriate for determining the conformational properties of the glyceryl side chains in the Neu5Ac residues, which determine the conformational properties of *NmB*. Our goal is to probe the effects of chemical modification on hapten conformation, and to provide a structural interpretation for the observed cross-reactivities or lack thereof between antibodies raised against the modified and native *NmB* CPS.

2. MATERIALS AND METHODS

2.1. Computational Methods. 2.1.1. MD Simulations. The strain B haptens (methyl α -(2,8)-di- and trisialosides) were built from the GLYCAM (32) library of monosaccharide residues using the XLEaP module of AMBER8 (34). The appropriate number of Na^+ counterions was added to neutralize the haptens using XLEaP, and the complete systems were immersed in pre-equilibrated boxes of 670 and 743 TIP3P (35) water molecules for the di- and trisialoside, respectively. Nonbonded van der Waals and electrostatic scaling factors for 1–4 interactions were set to unity (SCEE = SCNB = 1). Long range electrostatic interactions were computed with particle-mesh Ewald summation (36), with a nonbond cutoff distance of 8 Å. The constant pressure–temperature ensemble (nPT) was employed under periodic boundary conditions, with box dimensions of $30 \times 33 \times 35$ Å and $35 \times 31 \times 36$ Å for the di- and trisaccharide systems, respectively. Initially, the solvent configurations were energy minimized through 1000 cycles (50 steepest descent and 950 conjugate gradient minimization), while the CPS fragments were restrained. Subsequently, the entire system was energy minimized with the same protocol. Energy minimization was followed by a short simulated annealing stage, wherein the entire system was heated from 5 to 300 K over 50 ps, then cooled to 5 K over another 50 ps. Initial atomic velocities were assigned from a Boltzmann distribution at 5 K. Prior to the MD production cycle, the temperature of the system was raised to 300 K over 70 ps and held at that temperature for a further 80 ps. The production cycle ensued for 100 ns. A 2 fs time step was utilized to integrate the equations of motion, with hydrogen-containing bonds constrained to their equilibrium lengths using the SHAKE algorithm (37).

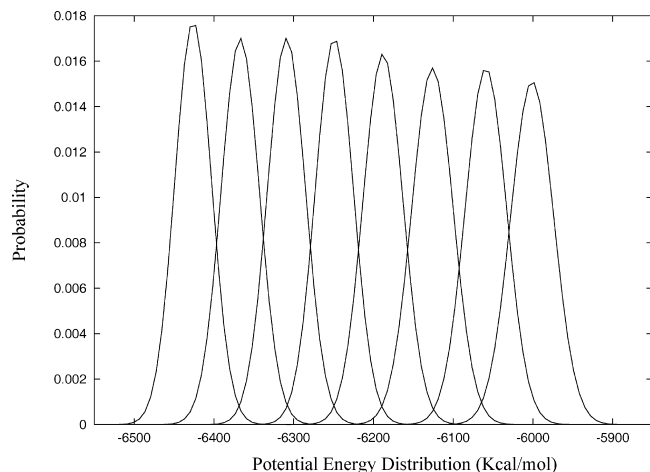


FIGURE 2: Gaussian potential energy distributions indicating the feasibility of the temperatures employed in the replica-exchange simulation.

2.1.2. Replica Exchange MD (REMD) Simulations. In REMD simulations, multiple replicas of a system are simulated concurrently at progressively higher target temperatures that increase exponentially (38, 39). At specified intervals, the energies of adjacent replicas are swapped based on an acceptance probability (39). Prior to performing the REMD simulations, the system was equilibrated following the same procedures outlined in the MD section. To derive the target temperatures employed in the present simulations, an exponential equation was fit to the range of sample temperatures given in the replica exchange section of the AMBER9 manual: 269.5, 300, 334, 371.8, 413.9, 460.7, 512.9 and 570.9 K. The exponential equation obtained from the fit was

$$y = 242.1 e^{(0.1072x)} \quad (1)$$

Eight approximate initial target temperatures (299.9, 308.0, 316.4, 324.9, 333.7, 342.8, 352.1 and 361.6 K) were generated by interpolating between the sample temperatures in incremental x -values of 0.25 starting from $x = 2.0$. From the initial single equilibration stage eight simulations were spawned by equilibrating to the eight approximate target temperatures noted above, followed by 10 ns of production dynamics under the constant volume–temperature (NVT) ensemble. Gaussian distributions (39) of the potential energies from these simulations were plotted, in order to confirm that energy overlaps occurred between successive replicas within the temperature range, Figure 2. Employing the lowest target temperature (300 K) from the simulations, and a swap acceptance probability of 0.2, the final target temperatures (300, 305, 310, 315, 320, 326, 331 and 337 K) were obtained through an iterative process described previously (39). Prior to committing the replicas to exchange dynamics, each of the eight systems was heated to its target temperature. The hybrid solvent REMD method (38) was employed retaining 100 closest water molecules during each replica exchange process. An exchange was attempted every 500th step for a total of 500×40000 steps, resulting in an overall simulation time of 8×20 ns for the eight replicas. A 1 fs time step was utilized to integrate the equations of motion, and a 10 kcal/mol restraint force constant was applied to the carbohydrate ring atoms to prevent the rings from distorting at high temperatures.

2.1.3. Bayesian Cluster Analysis. Cartesian coordinates from the MD simulations were collected every 5 ps for

subsequent analysis with either the CARNAL or PTRAJ modules of AMBER8. In addition to typical analysis of torsion angle distributions (30), global conformational states were also identified by determining transitions between Markov states, via a Bayesian statistical analysis of the glycosidic torsion angles (40). A 25-component mixture of univariate Gaussians was used as a fuzzy partition of the data space. The centers and the widths of these Gaussians were optimized by a maximum likelihood approach (41). Subsequently, the transition matrix for the 25 partition volumes was constructed. The two partition volumes connected by the fastest transitions were iteratively united. Thus, 25 Markov models and associated time scales of the fastest transitions between the states of the respective model were obtained. In this approach, if the time scale (τ_l) of model l is much larger than the time scale of the subsequent model $l + 1$ ($\tau_l \gg \tau_{l+1}$), then model $l + 1$ contains rapidly mixing states, while model l contains slowly varying metastable states. In that case, model l represents a plausible conformational state of the system. This enables the definition of several relevant state models for the observed trajectory (40), and identifies global relationships between glycosidic torsion angles that may not otherwise be readily determined.

2.1.4. Predicting NOE Intensities from the MD Trajectories of the Di- and Trisialosides. 2D NOE build-up intensities for the di- and trisialosides were predicted from MD simulations by performing a full relaxation matrix analysis with the CORMA program (42). A sample of 2500 snapshots, extracted at regular intervals from the MD trajectories, was utilized for the predictions. Correlation times of 0.5 and 0.55 ns were employed for the di- and trisialosides, respectively.

2.2. NMR Spectroscopy. Synthetic samples of the methyl α -2,8-di- and trisialosides were purchased from Dr. Nicolai Bovin (Lectinity Holdings, Lappeenranta, Finland). Samples were lyophilized from D_2O and then redissolved in 99.96% D_2O . Data were collected on Varian Inova 500, 800 and 900 MHz spectrometers at 25 °C, except for some NOE data which were also collected at 10 °C. All pulse programs were from the standard Varian library. Data were processed with Gaussian window functions, linear prediction, and zero-filling prior to Fourier transformation. For each sample, a COSY data set was collected with 1582×1024 complex points and acquisition times of 0.40 and 0.258 s in F2 and F1, respectively. An HSQC data set was collected with 789×400 complex points and acquisition times of 0.2 and 0.03 s in F2 and F1, respectively. Assignments are listed in Table 2. A second high-resolution HSQC spectrum was collected with the carbon decoupler turned off to allow a long acquisition time of 0.7 s during T2, in order to obtain proton–proton coupling data. 1D transient NOE experiments with mixing times of 100 to 800 ms were performed on the H3 axial signals in the Neu5Ac residues (H3ax) to determine the linear NOE buildup regime. Data was then collected, using a 300 ms mixing time, on those signals that had adequate spectral dispersion. Additional NOE data were obtained by acquiring a 2D NOESY spectrum utilizing a zero-quantum suppression element (43) and a mixing time of 300 ms. Data were collected with 420×2048 complex points and acquisition times of 0.459 and 0.09 s in F2 and F1. 1D data were processed with Varian VNMRJ software (Varian, Inc.), and 2D data were processed and analyzed

Table 2: Proton Chemical Shifts of Methyl α -(2,8)-Di- and Trisialoside, Acquired at 25 °C in D₂O, Relative to Internal DSS

	H3ax	H3eq	H4	H5	H6	H7	H8	H9	NAc	OMe
Methyl α -(2,8)-Disialoside										
b	1.565	2.612	3.552	3.783	3.813	3.858	4.194	4.133, 3.693	2.059	3.310
a	1.726	2.759	3.666	3.825	3.610	3.581	3.905	3.872, 3.635	2.022	
Methyl α -(2,8)-trisialoside										
c	1.572	2.607	3.531	3.801	3.838	3.890	4.144	4.141, 3.668	2.070	3.310
b	1.697	2.674	3.573	3.795	3.602	3.844	4.154	4.182, 3.674	2.062	
a	1.739	2.753	3.656	3.833	3.601	3.573	3.928	3.890, 3.634	2.022	

Table 3: Carbon Chemical Shifts of Methyl α -(2,8)-Di- and Trisialoside, Acquired at 25 °C in D₂O^a

	C3	C4	C5	C6	C7	C8	C9	NAc	OMe
Methyl α -(2,8)-Disialoside									
b	43.06	70.48	55.09	76.81	72.17	81.19	64.26	25.08	54.49
a	43.25	71.15	54.40	75.30	70.81	73.36	65.26	24.83	
Methyl α -(2,8)-Trisialoside									
c	42.95	70.67	55.19	76.37	71.87	80.71	64.10	25.16	54.54
b	42.88	70.88	55.09	76.47	72.08	81.10	64.18	25.09	
a	43.24	71.30	54.44	75.34	70.94	74.45	65.28	24.81	

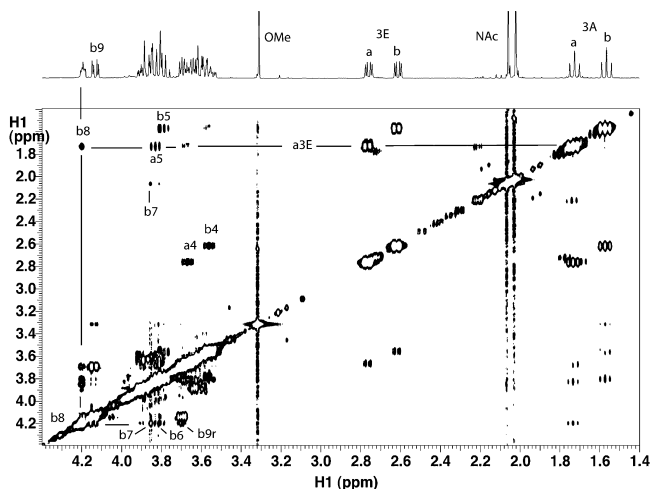
^a Referenced indirectly to DSS.

FIGURE 3: NOESY spectrum of disialoside collected at 10 °C and 800 MHz with a mixing time of 300 ms. A few cross-peaks used in determining the linkage conformation are indicated.

using NMRPipe software (44), while volume integrations were performed with the NMRView software (45).

2.2.1. Chemical Shift Assignments. Assignments were confirmed with COSY and HSQC data, Tables 2 and 3. Identification of the reducing, middle and terminal Neu5Ac residues was based on the existence of NOEs between the H3ax proton and the OMe aglycon, and between the H3ax/eq and H8 protons of the preceding residue, as well as in the relative downfield chemical shift of C8 due to the glycosidic linkage. *N*-Acetyl methyl groups were assigned based on NOEs to protons H5 and H7. Chemical shift data were consistent between **I** and **II** as can be seen by comparing the reducing and terminal residues of the di- and trisialosides, Tables 2 and 3.

2.2.2. NOE Measurements. The NOE data were collected at high field (800 and 900 MHz) and 10 °C to promote a stronger negative NOE. The 2D NOESY spectra of the disialosyl and trisialosyl haptens are shown in Figures 3 and 4, respectively. Due to the high degree of spectral overlap, only a limited number of NOEs could be confidently quantified from the many observed. A 1D selective NOESY

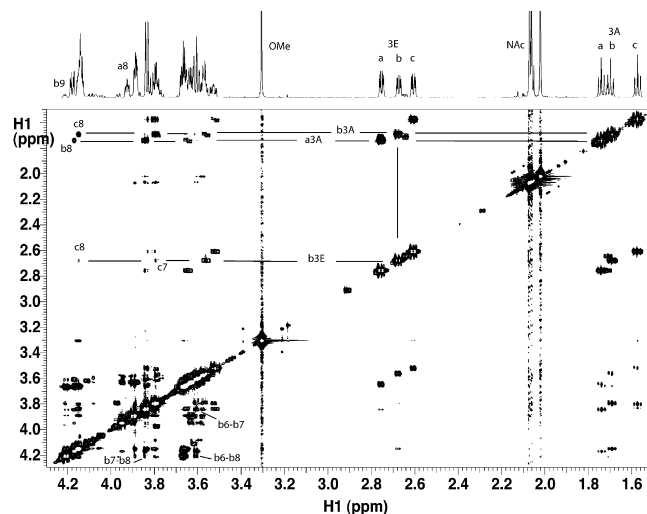


FIGURE 4: NOESY spectrum of trisialoside collected at 10 °C and 800 MHz with a mixing time of 300 ms. A few key cross-peaks used in determining the linkage conformation are indicated. Notably, weak cross-peaks between the equatorial H3 proton of the middle residue (bH3eq) and H7 and H8 of the reducing residue (cH7 and cH8) can be observed. Despite substantial overlap of signals, distinct cross-peaks can be measured for some ring protons (e.g., bH6–bH8) of the middle residue.

spectrum of the trisaccharide, Figure 5, clearly indicated a long-range NOE between protons H9 of the middle residue and H7 of the reducing residue. Peaks in 2D NOESY data sets, processed using NMRView, were integrated by defining boundary boxes around the peaks and summing points within the box according to the NMRView analysis function. 1D NOE data were integrated using standard Varian functions. Integrated values were used for distance calculations based on the isolated spin-pair approximation, wherein the H3eq to H3ax distance was set to 1.79 Å (9). Cross-peaks originating from the same source proton were then compared for their respective distances.

2.2.3. J-Coupling Measurements. Extracting accurate scalar coupling constants from strongly coupled protons frequently found in carbohydrates can be achieved by a combination of selective 1D TOCSY spectra, followed by a

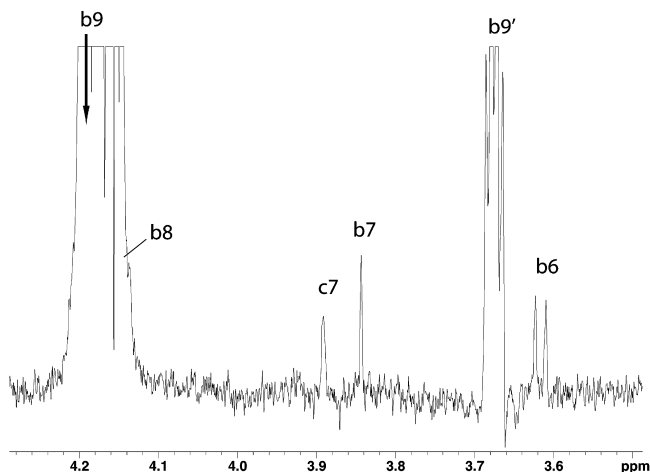


FIGURE 5: The 1D NOE of trisialoside. Expected intraresidue NOEs from proton H9 of the middle residue (bH9) to adjacent protons (bH6, bH7, bH8, bH9') are shown, in addition to a long-range NOE to the H7 of the reducing residue (cH7).

simulation of the resulting spin system. In the *NmB* haptens, selective TOCSY methods were limited by the poor transfer from protons H6 to H7, and from overlapping resonances that prevented access to the important H7, H8 and H9 protons. Thus a 2D HSQC spectrum was acquired with high digital resolution in the proton dimension. The high-resolution HSQC data were Fourier transformed with additional linear prediction and zero-filling in T2 and T1 in order to maximize digital resolution to 1.0 Hz per point. Traces through the carbon-proton peaks provided, in most cases, proton subspectra showing first-order coupling. J -values were measured directly from these traces.

3. RESULTS AND DISCUSSION

3.1. NMR and Conformational Properties. The conformational properties of the α -(2,8)-linked Neu5Ac residues are largely determined by the rotatable bonds between the pyranose rings (9, 46). Previous MD simulations of **I** indicated some pyranose ring interconversions from the predominant 2C_5 chair conformation (29), which were rationalized only via nonaqueous phase experimental and molecular mechanics studies. However, there were no indications that these ring fluctuations influenced the conformational properties of the dimer (29). The predominant ring configuration was determined from the scalar ${}^3J_{HH}$ -values of the ring protons. In the 2C_5 conformation the dihedral angles between protons H3ax–H4, H4–H5 and H5–H6 are *anti*, while that between H3eq–H4 is *gauche*. These rotamer orientations give rise to high and low magnitude 3J -values for protons in the *anti*- and *gauche*-orientations, respectively. The theoretical scalar J -coupling constants were computed employing a Karplus equation that takes into account the electronegativities and orientations of substituent groups (47). For example, in residue (a) of the disialoside, values for *anti*-orientations of 11.7 ± 0.7 Hz, 9.4 ± 0.8 Hz and 10.4 ± 0.5 Hz for the H3ax–H4, H4–H5 and H5–H6 protons, respectively, computed from the MD simulation were in good agreement with the experimental values of 12.2 ± 0.1 Hz, 10.2 ± 0.1 Hz and 10.2 ± 0.1 Hz, respectively. A value of 4.6 ± 1.2 Hz was computed for the *gauche* H3eq–H4 proton pair from the MD data, which also showed good agreement

Table 4: NMR and Computed ${}^3J_{HH}$ Coupling Constants (Hz) for Inter-Residue Torsion Angles in Fragments of *NmB* Sialobioside, α -Neu5Ac-(2-8)- α -Neu5Ac-OMe (a–b)

angle	linkage	spins	NMR	MD (100 ns) ^a	REMD (20 ns) ^{a,b}
ω_7	terminal	aH6–aH7	1.4 ± 0.1	1.0 ± 0.8	1.0 ± 0.8
	internal	bH6–bH7	<1.0	1.2 ± 0.9	1.2 ± 0.9
ω_8	terminal	aH7–aH8	9.5 ± 0.9	7.9 ± 0.6	8.1 ± 0.6
	internal	bH7–bH8	1.5 ± 0.2	2.0 ± 0.7	2.0 ± 0.7
ω_9	terminal	aH8–aH9R	6.1 ± 0.6	6.7 ± 1.5	5.9 ± 1.5
	(free rotor)				
		aH8–aH9S	2.4 ± 0.2	3.7 ± 0.9	3.6 ± 0.9
	internal	bH8–bH9R	6.1 ± 0.6	7.3 ± 1.6	7.4 ± 1.7
	(free rotor)				
		bH8–bH9S	4.1 ± 0.4	2.9 ± 1.3	2.9 ± 1.3

^a Conformation-defining intraresidue J -couplings are shown in bold. ^b REMD simulations were performed at 8 temperatures for a total of 160 ns.

with the experimental value of 4.4 ± 0.4 Hz. A complete list of the pyranosyl ${}^3J_{HH}$ -values is presented in Table S1 in the Supporting Information. These J -values indicated that the pyranosyl rings remained entirely in the 2C_5 conformation although the ring torsion angles displayed modest oscillations of approximately $\pm 15^\circ$. Therefore, the conformational properties of the methyl α -(2,8)-di- and trisialosides were characterized here by monitoring the time-dependent evolution or populations of the φ , ψ , ω_8 and ω_7 angles (see Figure 1) during the MD and REMD simulations.

In order to ascertain whether the results from the present simulations were consistent with experiment, and so could be relied on to shed some light on the conformational properties of the strain B haptens, interproton distances and scalar ${}^3J_{HH}$ -coupling constants were computed from the MD trajectories and compared with experimental data determined herein.

3.1.1. Methyl α -(2,8)-Disialoside (a–b-OMe). The experimental NMR coupling constants and those computed from the MD simulations of **I** are presented in Table 4. It can be seen that the values derived from the MD and REMD simulations were statistically indistinguishable, and the back-calculated ${}^3J_{HH}$ values showed good overall agreement with the experimental coupling constants. In terms of conformational properties, the small experimental ${}^3J_{bH6-bH7}$ and ${}^3J_{bH7-bH8}$ values of <1.0 and 1.5 Hz, respectively, for these sets of protons in the internal reducing Neu5Ac residue, were consistent with a \pm *gauche* orientation, due to the symmetry of the Karplus relationship (47). The precise rotamer assignment required the utilization of NOE data, Table 5. The strong NOEs measured between the bH8–bH6 and bH8–bH7 protons confirmed the $-gauche$ and $+gauche$ conformations for the bH6–bC6–bC7–bH7 and bH7–bC7–bC8–bH8 dihedral angles, respectively, which corresponded to a $+gauche$ orientation for both the ω_7 (bO6–bC6–bC7–bO7) and ω_8 (bO7–bC7–bC8–bO8) angles. In the nonreducing terminal residue, the small magnitude of the ${}^3J_{aH6-aH7}$ coupling constant (1.4 Hz) for the glyceryl side chain indicated a $+gauche$ conformation for the ω_7 (aO6–aC6–aC7–aO7) angle. In contrast, the large ${}^3J_{aH7-aH8}$ value of 9.5 Hz for this residue suggested an *anti* orientation for these protons, which corresponded to an *anti* orientation for this ω_8 (aO7–aC7–aC8–aO8) torsion angle. These results are similar to those reported previously and indicate that the orientation of ω_8 depends heavily on whether O8 is glycosylated or free (9, 46). The MD and REMD simulations gave

Table 5: Experimental (ISPA) NOE Distances (\AA) and Average Interproton Distances from MD and REMD Simulations for Fragments of NmB Sialobioside, α -Neu5Ac-(2-8)- α -Neu5Ac-OMe (a-b)^a

spins	NMR	MD (100 ns)	REMD (20 ns)	spins	NMR	MD (100 ns) ^b	REMD (20 ns) ^{b,c}
aH3ax-aH3eq	1.8 ± 0.02	1.7 ± 0.1	1.7 ± 0.1	bH8-bH6	2.9 ± 0.03	2.5 ± 0.2	2.5 ± 0.2
aH3ax-aH4	3.4 ± 0.04	3.0 ± 0.1	3.0 ± 0.1	bH8-bH7	2.5 ± 0.03	2.6 ± 0.2	2.6 ± 0.2
aH3ax-aH5	2.8 ± 0.03	2.6 ± 0.2	2.7 ± 0.2	aH3ax-bH8	3.0 ± 0.03	2.5 ± 0.8	2.5 ± 0.9
aH3eq-aH4	2.8 ± 0.03	2.4 ± 0.1	2.4 ± 0.1	aH3eq-bH8	4.0 ± 0.04	3.8 ± 0.5	3.8 ± 0.5
bH3ax-bH4	3.2 ± 0.04	3.0 ± 0.1	3.0 ± 0.1	bH5-bNMe	4.3 ± 0.05	4.5 ± 0.1	4.5 ± 0.1
bH3ax-bH5	3.0 ± 0.04	2.6 ± 0.2	2.6 ± 0.2	bH9R-bH8	2.5 ± 0.03	2.8 ± 0.3	2.8 ± 0.3

^a The distances are averaged over the MD trajectories as $\langle 1/R^6 \rangle^{(-1/6)}$. ^b Conformation-defining intra- and inter-residue NOE's are shown in bold. ^c REMD simulations were performed at 8 temperatures for a total of 160 ns.

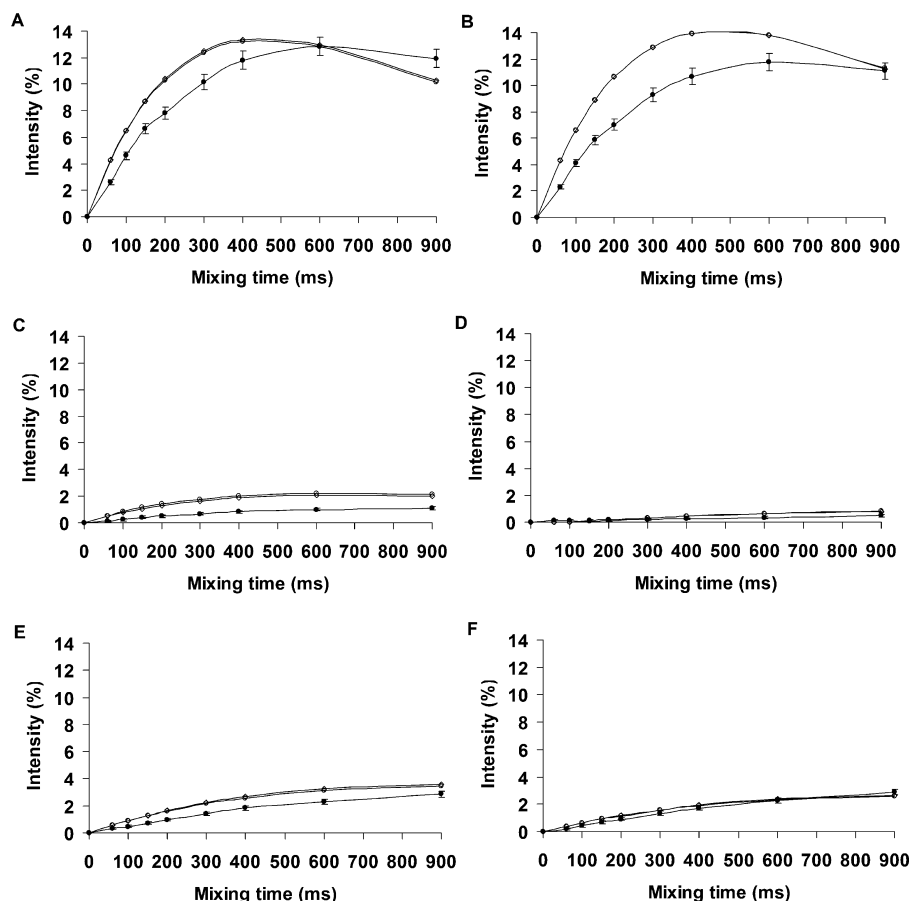


FIGURE 6: Experimental and theoretical NOE build-up curves of proton pairs in disialoside (a-b-OMe). A, aH3ax-aH3eq; B, bH3ax-bH3eq; C, aH3ax-bH8; D, aH3eq-bH8; E, bH8-bH6; F, bH8-bH7. A correlation time of 0.5 ns was employed in the theoretical predictions. Experiment \bullet ; conventional MD \circ ; REMD \diamond .

similar NOE values, which compared well with the experimentally derived distances, Table 5. The interglycosidic NOE distances were largely determined by the linkage dihedral angles; therefore the fact that the MD simulations were sampling experimentally consistent torsion space was also supported by the good agreement between the experimental and the MD-derived interglycosidic NOE distances. For example the NOE distances measured between the aH3ax-bH8 protons was $2.5 \pm 0.8 \text{ \AA}$ for both the MD and REMD, respectively, which may be compared with the experimental value of $3.0 \pm 0.03 \text{ \AA}$. The experimentally derived NOE distance between the aH3eq-bH8 protons was $4.0 \pm 0.04 \text{ \AA}$, compared to $3.8 \pm 0.5 \text{ \AA}$ for both the MD and REMD simulations.

To determine the fit between the experimental and theoretical NOE intensities in **I**, NOE build-up curves were computed employing a complete relaxation matrix analysis. A single overall correlation time of 0.5 ns for all pairs of

protons resulted in good agreement between the experimental and theoretical data, Figure 6. Notably, the theoretical data gave rise to NOE intensities that were in quantitative agreement was excellent for the conformationally significant proton pairs (aH3ax-bH8, aH3eq-bH8, bH8-bH6 and bH8-bH7) over the entire range of mixing times.

The conformational itineraries of the glyceryl exocyclic dihedral angles from the 100 ns MD simulation of **I** are presented in Figure 7. Despite the potential for the glycosidic linkage to populate a wide range of conformations, the data from the MD and REMD simulations suggested that well-defined subsets of conformations were preferentially adopted. The ω_7 angle predominantly sampled the +gauche rotamer in both Neu5Ac residues. In contrast, the ω_8 angle exhibited a preference for several staggered rotamers depending on whether the O8 atom was involved in a glycosidic bond. Consistent with the experimental vicinal $^3J_{H7-H8}$ coupling constants, the ω_8 angle was predicted to exist predominantly

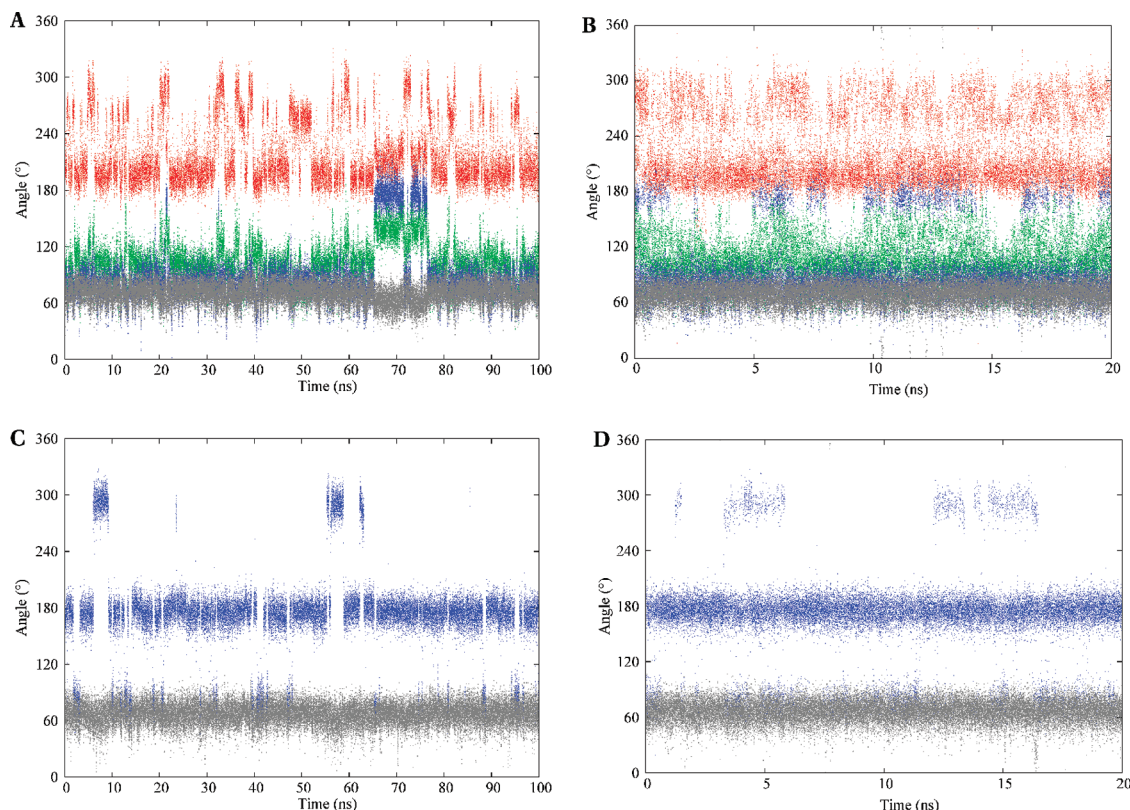


FIGURE 7: The conformational variations of the exocyclic glyceryl dihedral angles of the methyl α -(2,8)-disialoside. ϕ , red; ψ , green; ω_8 , blue; ω_7 , gray. The A–B and C–D panels depict dihedral angle variations of the glyceryl region involved and not involved in glycosidic bond formation, respectively. The left and right columns represent trajectories from the conventional MD and REMD simulations, respectively.

in the *anti* orientation when the O8 atom was not involved in a glycosidic bond, but preferred the *gauche* orientation otherwise. The predominance of the *anti* conformation when O8 is nonglycosidically linked has been observed in X-ray crystal structures (48) and inferred from solution-phase NMR (9, 46, 49) data. Two intraresidue hydrogen bonds, between the oxygen atoms of the carboxylate group with the hydroxyl group at C8, and between the carbonyl group of the acetamido functionality and the hydroxyl group at C7, seem to be the origin of this rotamer's stability (49). However, the magnitude of this stabilization is not clear, since a similar *anti* orientation for the ω_8 angle has been observed in the X-ray crystal structures of the β -anomer (48), in which the formation of the intraresidue hydrogen bond to the carboxylate group is impossible.

The ϕ -angle in Neu5Ac residues is known to populate two conformations, *-gauche* and *anti* (49, 50). There are no conclusive experimental data on the solution conformational properties of the ψ angle; however, potential energy calculations suggest that the ψ angle oscillates between 105° and 130° (9). Over the course of the MD simulations, correlated motions were observed between the conformational itineraries of the ω_8 , and ϕ and ψ dihedral angles of the glycosidic linkage region. During the first 65 ns, the ω_8 angle predominantly populated the *+gauche* rotamer ($78 \pm 13^\circ$), the ϕ angle displayed transitions between its two solution preferred conformations, ($-161 \pm 14^\circ$, $-91 \pm 17^\circ$), while the ψ angle adopted a value of $98 \pm 18^\circ$. After 65 ns the ω_8 angle effected a transition to the *anti* rotamer. During the time interval that the ω_8 -*anti* rotamer persisted, if the ϕ angle adopted either the *-gauche* ($-91 \pm 17^\circ$) or *anti* ($-161 \pm$

14°) conformations then a *+gauche* ψ ($98 \pm 18^\circ$) would introduce unfavorable electrostatic repulsions due to the close proximity of the carboxyl groups and ring oxygen atoms on two adjacent Neu5Ac residues, Figure 8. Presumably to alleviate these interactions, the ψ angle flips to the *anti* rotamer ($142 \pm 12^\circ$). In general, the preferred conformational preferences of the exocyclic dihedral angles observed during the MD simulations are in accordance with previous reports from experimental NMR studies of the respective angles (9, 46, 50). The majority of the transitional correlations between the ω_8 , and ϕ and ψ angles were also observed in the REMD simulation. Comparisons of the relative distributions of the rotamers of the exocyclic glyceryl side chain computed from experimental *J*-values, as well as the MD and REMD simulations, are shown in Table 6. For the conformationally significant dihedral angles, particularly ω_7 and ω_8 , the theoretical populations showed good agreement with experiment. Only a qualitative agreement was observed between the theoretical and experimental populations for the freely rotating ω_9 angle.

3.1.2. Methyl α -(2,8)-Trisialoside (*a-b-c-OMe*). Methyl α -(2,8)-trisialoside (**II**) was included in this study to investigate the effects of multiple Neu5Ac residues on the conformational properties of the 2,8-linkage. To the best of our knowledge only two experimental studies have been carried out on the trisaccharide hapten (9, 46). In those studies the trisaccharide fragment terminated in the β -configuration, limiting the relevance of those data. The vicinal homonuclear $^3J_{\text{HH}}$ -coupling constants in **II**, determined experimentally and those from our MD simulations, are presented in Table 7. The observed $^3J_{\text{H6-H7}}$ constants for all

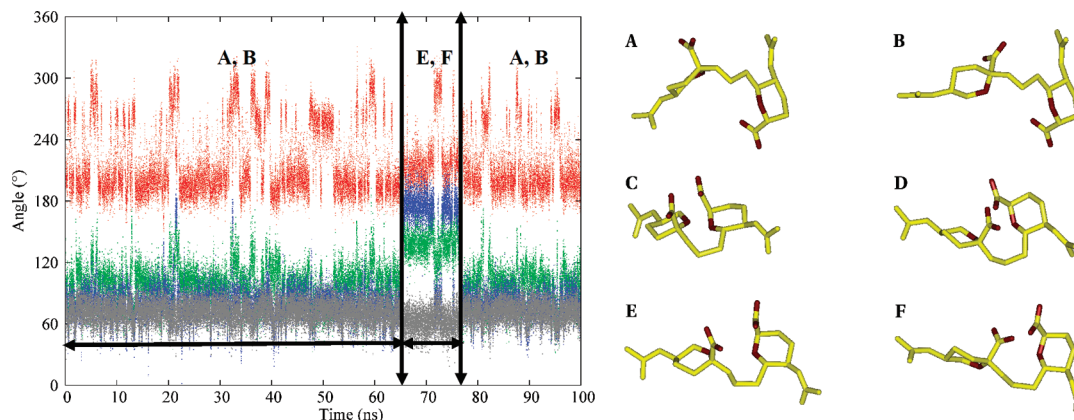


FIGURE 8: Possible repulsive interactions between the charged carboxyl groups and the O6 ring oxygen atoms on two adjacent residues. $(\varphi, \psi, \omega_8) = (-161, 98, 78)$ and $(-91, 98, 78)$ in A and B, respectively, denoting the solution-preferred structures during the first 65 ns of the simulation. $(\varphi, \psi, \omega_8) = (-161, 98, 175)$ and $(-91, 98, 175)$ in C and D, respectively, represent unfavorable interactions that would occur if the ω_8 angle adopted the *anti* rotamer while the ψ angle remained in the +*gauche* (98°) state. To alleviate the repulsive interactions, the ψ angle makes a transition to the *anti* rotamer, E and F with $(\varphi, \psi, \omega_8) = (-161, 142, 175)$ and $(-91, 142, 175)$, respectively.

Table 6: NMR-Based and Computed Populations (%) for Inter-Residue Torsion Angles in Fragments of *NmB* Disialoside, α -Neu5Ac-(2-8)- α -Neu5Ac-OMe (a–b)

angle	linkage	NMR ^{a,b,c}	MD (100 ns) ^a	REMD (20 ns) ^{a,e}
ω_7	terminal a	100/0/0	100/0/0	100/0/0
ω_7	internal b	100/0/0	100/0/0	100/0/0
ω_8	terminal a	0/100 ^d /0	13/80/7	13/83/4
ω_8	internal b	96 \pm 3/3 \pm 3/0	90/10/0	90/10/0
ω_9	terminal a	54 \pm 8/0/50 \pm 12	61/18/21	52/16/32
ω_9	internal b	55 \pm 8/24 \pm 9/20 \pm 15	73/2/25	75/4/21

^a *gauche/trans/–gauche*. ^b Range is consistent with the estimated error in the experimental *J*-values, only one symmetric solution is presented. ^c NMR populations derived by employing limiting *J*-values from rotational isomeric states computed from conventional MD (the REMD simulations did not give significantly different limiting *J*-values). ^d Estimated based on the magnitude of the *J*-value. ^e REMD simulations were performed at 8 temperatures for a total of 160 ns.

Table 7: Conformationally Sensitive NMR and Computed $^3J_{\text{HH}}$ Coupling Constants (Hz) for Inter-Residue Torsion Angles in Fragments of *NmB* Trisialoside, α -Neu5Ac-(2-8)- α -Neu5Ac-(2-8)- α -Neu5Ac-OMe (a–b–c)

angle	linkage	spins	NMR	MD (100 ns) ^a	REMD (20 ns) ^a
ω_7	terminal	aH6–aH7	1.5 \pm 0.2	1.0 \pm 0.8	1.0 \pm 0.8
	internal	bH6–bH7	<1.0	0.9 \pm 0.8	0.9 \pm 0.7
	internal	cH6–cH7	<1.0	1.1 \pm 0.9	1.3 \pm 1.0
ω_8	terminal	aH7–aH8	9.6 \pm 1.0	7.7 \pm 0.6	8.1 \pm 0.5
	internal	bH7–bH8	<4.0	3.6 \pm 0.9	1.9 \pm 0.9
	internal	cH7–cH8	<4.0	2.1 \pm 0.7	2.1 \pm 0.8

^a REMD simulations were performed at 8 temperatures for a total of 160 ns.

the Neu5Ac residues from the experiment were less than 1.5 Hz, which indicated a *gauche* relationship between protons H6 and H7. The average $^3J_{\text{H6–H7}}$ values computed from the MD-simulation were also of low magnitude, and were in excellent agreement with the experimental values. As in the case of the disaccharide, the large experimental $^3J_{\text{aH7–aH8}}$ value of 9.6 Hz indicates that these protons are predominantly in the *anti* orientation for the nonreducing terminal residue. The MD-computed value for $^3J_{\text{aH7–aH8}}$ (7.7 \pm 0.6 Hz) also suggested a predominantly *anti* orientation, which upon further analysis was confirmed to be 78% *anti* and 22% *gauche*. In contrast to the large $^3J_{\text{aH7–aH8}}$ values observed in the nonreducing terminal residue, the $^3J_{\text{H7–H8}}$ values for the

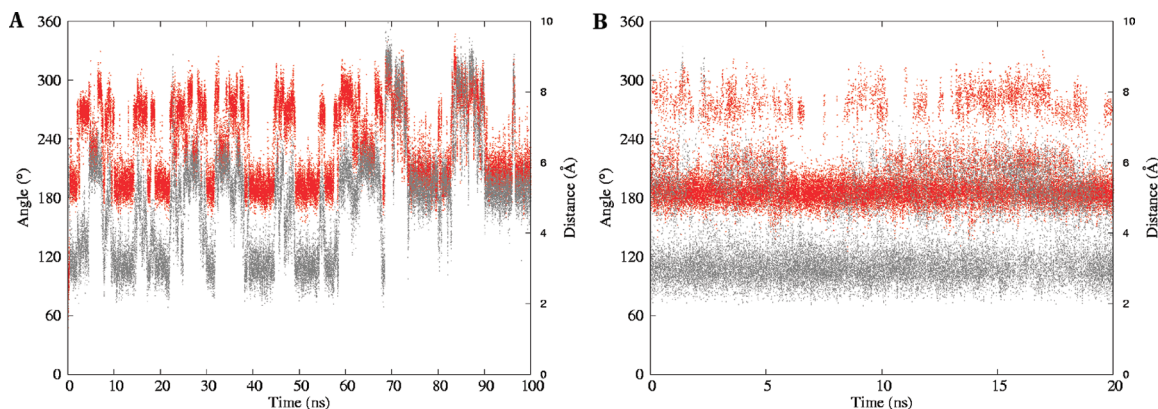
internal (b) and reducing-internal (c) residues, in which O8 is involved in glycosidic bonds, were less than 4.0 Hz, indicating a conformational change to the *gauche* orientation. That a *gauche* rotamer was preferred for the a–b and b–c linkages, respectively, was also demonstrated by the relatively small magnitude of the $^3J_{\text{H7–H8}}$ coupling constants observed in the MD data (2.1 \pm 0.7 and 2.1 \pm 0.8 for the MD and REMD simulations, respectively). In addition, the small magnitude of the $^3J_{\text{H7–H8}}$ values for the residues b and c implies similarities between the conformational properties of the ω_8 angle in both the a–b and b–c linkages. In previous experimental NMR studies of the Neu5Ac trisaccharide (9, 46) the anomeric center of the reducing-terminal Neu5Ac residues was in the β -configuration, which led to the conclusion that the conformational properties of the ω_8 angle in the two linkage regions differed (46). Here, for the all α -configuration, we see that this is not the case. Good agreement was also obtained between the experimental and MD-computed values for the $^3J_{\text{H8–H9R}}$ and $^3J_{\text{H8–H9S}}$ couplings; however, the rotameric preferences of the C8–C9 bond are not directly pertinent to the overall 3D conformational properties of the CPS.

NOE distances were computed from the MD and REMD simulations of the trimer and compared with the available experimental data, Table 8. Unlike the case of the dimer, the agreement with the experimental values in the trisaccharide was only qualitative, suggesting a need for longer sampling time. The NOE values from the REMD simulation were in better agreement relative to the MD simulation, except for one case, bH3eq–cH7. It should be noted that a very weak ($>1\%$) long-range NOE was observed between protons H9 of the internal residue (b) and H7 of the reducing terminal residue (c), Figure 5. The ISPA-derived experimental distance between these protons was 3.1 \pm 0.06 Å. Given that the bH9 protons were not unambiguously identified, the center of mass of the bH9R and bH9S protons was employed to determine the bH9–cH7 distance from the MD trajectories. Interproton distances of 3.0 \pm 1.4 and 3.5 \pm 2.1 Å were obtained from the REMD and MD data, respectively. Further investigations revealed that the bH9–cH7 distance was highly sensitive to the φ -angle of the b–c linkage, Figure 9. In particular, these protons were within

Table 8: Experimental (ISPA) NOE Distances (Å) and Average Interproton Distances from MD and REMD Simulations for Fragments of NmB Trisialoside, α -Neu5Ac-(2-8)- α -Neu5Ac-(2-8)- α -Neu5Ac-OMe (a-b-c)^a

spins	NMR	MD (100 ns) ^b	REMD (20 ns) ^{b,c}	spins	NMR	MD (100 ns) ^b	REMD (20 ns) ^{b,c}
bH3ax-cH8	2.6 ± 0.06	2.6 ± 1.0	2.3 ± 0.8	bH6-bH8	2.3 ± 0.05	2.5 ± 0.4	2.4 ± 0.2
bH3eq-cH8	3.5 ± 0.08	3.9 ± 0.5	3.6 ± 0.5	aH3ax-bH8	2.8 ± 0.07	2.7 ± 1.1	2.3 ± 0.8
bH3eq-cH7	3.9 ± 0.09	4.6 ± 0.4	5.0 ± 0.5	aH3eq-bH8	3.4 ± 0.08	4.0 ± 0.5	3.6 ± 0.4
bH6-bH7	2.2 ± 0.05	2.4 ± 0.1	2.4 ± 0.1	bH9-cH7 ^d	3.1 ± 0.06	3.5 ± 2.1	3.0 ± 1.4

^a The distances are averaged over the MD trajectories as $\langle 1/R^6 \rangle^{-1/6}$. ^b Conformationally insensitive NOEs for the trisialoside were similar to the disialoside values and are omitted. ^c REMD simulations were performed at 8 temperatures for a total of 160 ns. ^d Given that the bH9 protons were not unambiguously identified, the center of mass of the bH9R and bH9S protons was employed to determine the bH9-cH7 distance from the MD trajectories.

FIGURE 9: Correlation between the bH9-cH7 interproton distance (gray) and the rotameric properties of the ϕ -angle (red) of the b-c linkage in the trisaccharide (a-b-c-OMe). Traditional MD (A); replica exchange MD (B).Table 9: NMR-Based and Computed Populations (%) for Inter-Residue Torsion Angles in Fragments of NmB Trisialoside, α -Neu5Ac-(2-8)- α -Neu5Ac-(2-8)- α -Neu5Ac-OMe (a-b-c)

angle	linkage	NMR ^{a,b,c}	MD (100 ns) ^a	REMD (20 ns) ^{a,e}
ω_7	terminal a	100/0/0	100/0/0	100/0/0
ω_7	internal b	100/0/0	100/0/0	100/0/0
ω_7	internal c	100/0/0	100/0/0	100/0/0
ω_8	terminal a	0/100 ^d /0	15/74/7	14/82/4
ω_8	internal b	(60 ± 9)/ (25 ± 3)/0	70/30/0	92/8/0
ω_8	internal c	(66 ± 5)/ (31 ± 9)/0	89/11/0	82/10/8

^a *gauche/trans/-gauche*. ^b Range is consistent with the estimated error in the experimental *J*-values, only one symmetric solution is presented. ^c NMR populations derived by employing limiting *J*-values from rotational isomeric states computed from conventional MD (the REMD simulations did not give significantly different limiting *J*-values). ^d Estimated based on the magnitude of the *J*-value. ^e REMD simulations were performed at 8 temperatures for a total of 160 ns.

the range of a detectable NOE (<3.5 Å) only when the ϕ -angle adopted the *anti* rotamer. Since the MD data displayed a mixture of both the *anti* and *gauche* rotamers for this angle, the reproduction of the bH9-cH7 distance can be utilized as an indicator of whether a simulation is correctly sampling the ϕ -angle rotamers in polysialic acids. The average distances between protons bH9 and cH7, computed from the first 70 ns of the conventional MD simulation for the periods when the ϕ -angle is *anti* (52%) or *-gauche* (48%), were 3.3 ± 1.8 Å and 5.3 ± 1.5 Å, respectively. Using these values the experimental distance of 3.1 Å equates to a ratio of *anti*:-*gauche* rotamers of approximately 100:0. The MD simulation therefore underestimates the population of the *anti* rotamer, leading to a slightly long value for the bH9-cH7 distance. Despite the marked improvement of the NOE-consistent population during the REMD simulation, these observations suggest a

possible need for improvement in the force field parametrization for the ϕ -angle in Neu5Ac. Nonetheless, the better performance of the REMD simulation indicates that enhanced-sampling methods can be effectively employed in studying the conformational properties of larger fragments of polysaccharides. Theoretical and experimental NOE build-up intensities in **II**, computed with the utilization of a complete relaxation matrix, are shown in Figure 10. The subtle variations in the bH9-cH7 NOE notwithstanding, a correlation time of 0.55 ns resulted in excellent overall agreement between the experimental and theoretical data for all pairs of protons. Rotamer populations computed from the experimental *J*-values and directly from the MD simulations are shown in Table 9. In general, the agreement with experiment was excellent with the exception of the ω_8 populations of the internal linkages, for which the theoretical data, nevertheless, correctly ranked the rotameric preferences.

The time series for the rotational angles of the glyceryl side chain in **II** over the 100 ns simulation are presented in Figure 11. In all three Neu5Ac residues the ω_7 angle populated only the +*gauche* rotamer. The ω_8 angle predominantly adopted the *anti* conformation in the glyceryl side chain of the nonreducing terminal residue, while in the internal and reducing-terminal residues the +*gauche* rotamer was preferred. Over the course of the simulation, the ψ angle oscillated around $101 \pm 30^\circ$ and $109 \pm 27^\circ$ for the a-b and b-c linkages, respectively, while the ϕ angle frequently populated the *anti* and *-gauche* rotamers in both linkage regions. As in the case of the linkage properties of **I**, correlated motions were observed between the ω_8 , and ϕ and ψ dihedral angles, which were strongly evident for the terminal (a-b) linkage in **II**. During the first 70 ns, the ω_8 angle predominantly populated the +*gauche* ($76 \pm 12^\circ$) rotamer and the ϕ angle displayed transitions between its

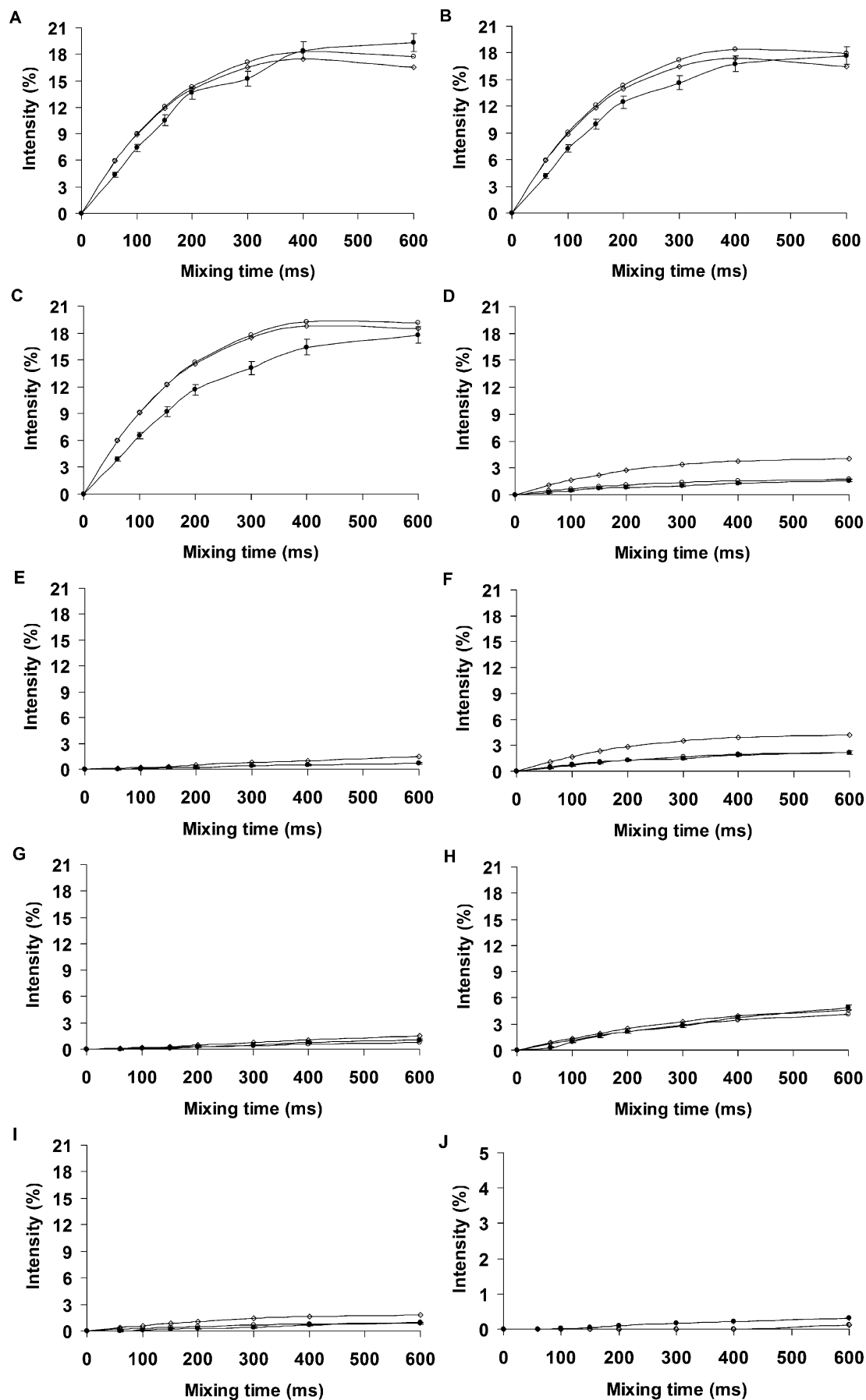


FIGURE 10: Experimental and theoretical NOE build-up curves for proton-pairs in the trisaccharide. A, aH3ax–aH3eq; B, bH3ax–bH3eq; C, cH3ax–cH83eq; D, aH3ax–bH8; E, aH3eq–bH8; F, bH3ax–cH8; G, bH3eq–cH8; H, bH8–bH6; I, bH9–cH7; J, bH3e–cH7. A correlation time of 0.55 ns was employed in predicting the theoretical NOE intensities. Experiment ●; conventional MD ○; REMD ◇.

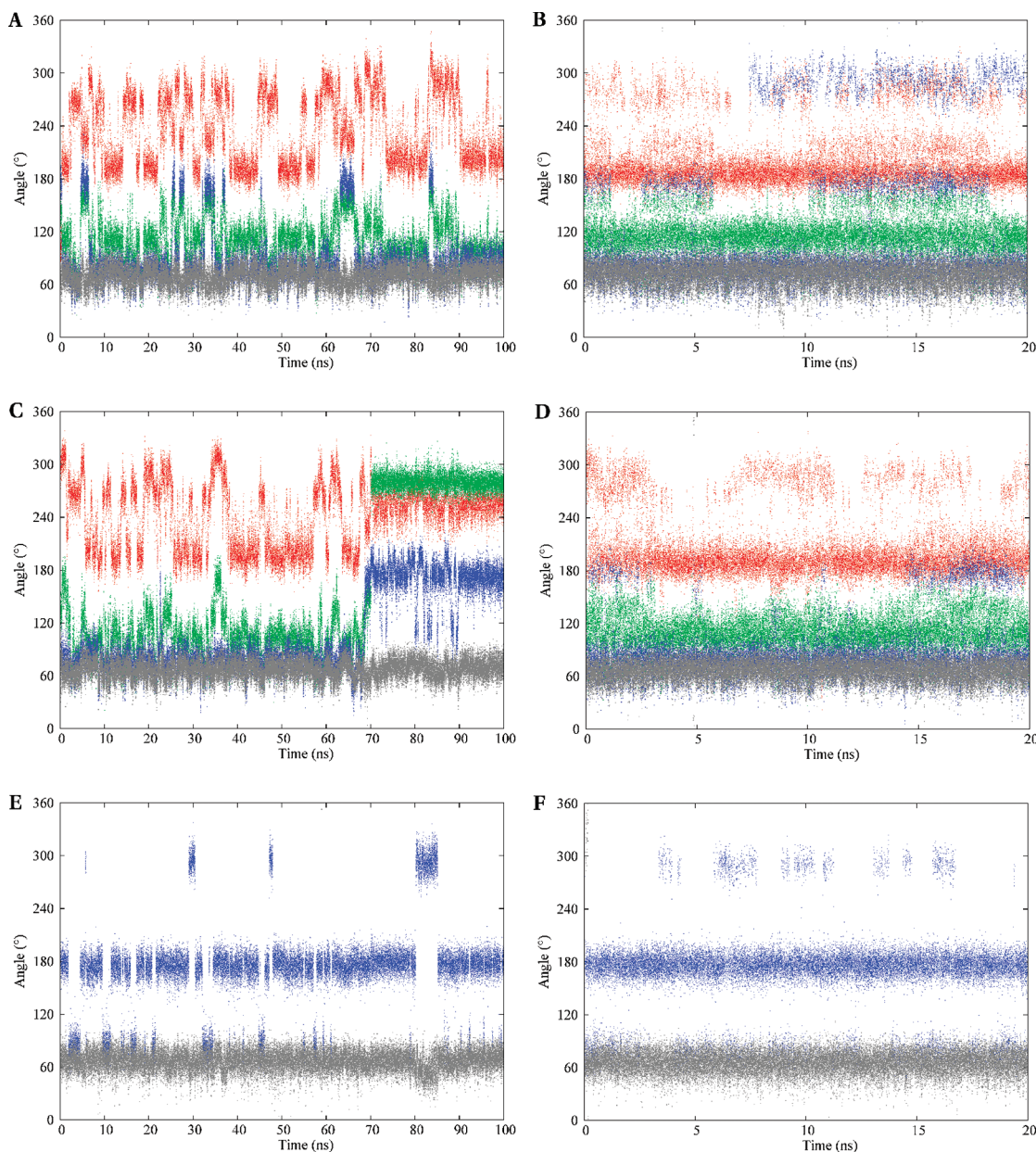


FIGURE 11: The conformational variations of the exocyclic glyceryl dihedral angles of the methyl α -(2,8)-trisialoside. φ , red; ψ , green; ω_8 , blue; ω_7 , gray. The top, middle and bottom panels represent trajectories from the reducing, internal, and terminal glyceryl side chains, respectively. The left and right columns represent the conventional MD and REMD simulations, respectively.

two solution preferred conformations ($-159 \pm 14^\circ$ and $-84 \pm 19^\circ$). After 70 ns the ω_8 dihedral angle effected a transition to the *anti* ($174 \pm 15^\circ$) rotamer. The presumed unfavorable electrostatic repulsions, discussed earlier in the disaccharide, again resulted in the ψ and φ angles effecting conformational transitions to their respective *-gauche* rotamers, $-80 \pm 8^\circ$ and $-104 \pm 14^\circ$, respectively. However, unlike in the case of the disaccharide, these rotamers persisted for the remainder of the simulation. Despite the plausible explanation for the existence of these rotamers observed during the last 30 ns of the conventional MD simulation, we sought to determine whether they represented experimentally relevant conformations.

Therefore, J -values and interproton distances were computed pre- and postconformational transition of the ω_8 angle, the first 70 and last 30 ns, respectively. A closer observation of the J -values computed from the conventional MD and REMD simulations indicated that the J -values were statistically similar except for the $^3J_{\text{bH7-bH8}}$ values, where the φ , ψ

and ω_8 transitions occurred. The value computed from the conventional MD simulation was about a factor of 2 (3.6 ± 0.9 Hz) greater than that determined from the REMD simulation (1.9 ± 0.9 Hz). The higher value may be due to contributions from the *anti* rotamers of ω_8 . The J -values computed before and after the conformational transition are presented in Table 10. These values are statistically similar and in good agreement with the experimental data except the $^3J_{\text{bH7-bH8}}$ values that are 1.4 ± 0.8 Hz and 8.7 ± 0.1 Hz for the first 70 and last 30 ns, respectively. The large value of 8.7 ± 0.1 Hz suggests that the last 30 ns may not be entirely experimentally relevant. However, the value of 1.4 ± 0.8 Hz is still small relative to the experimental value of <4.0 Hz, suggesting that the conformational properties of the ω_8 angle may be described by contributions from both the *+gauche* and *anti* rotamers, with a predominance of the *+gauche* rotamer.

Table 10: NMR and Computed $^3J_{\text{HH}}$ Coupling Constants (Hz) for Inter-Residue Torsion Angles in Fragments of *NmB* Trisialoside during the First 70 ns and Last 30 ns of the Conventional MD Simulation

angle	linkage	spins	NMR	MD
Preconformational Transition (The First 70 ns)				
ω_7	terminal	aH6–aH7	1.5 ± 0.2	1.0 ± 0.8
	internal	bH6–bH7	<1.0	0.9 ± 0.7
	internal	cH6–cH7	<1.0	1.1 ± 0.9
ω_8	terminal	aH7–aH8	9.6 ± 1.0	7.6 ± 0.6
	internal	bH7–bH8	<4.0	1.4 ± 0.8
	internal	cH7–cH8	<4.0	2.4 ± 0.7
Postconformational Transition (The Last 30 ns)				
ω_7	terminal	aH6–aH7	1.5 ± 0.2	1.0 ± 0.8
	internal	bH6–bH7	<1.0	1.0 ± 0.8
	internal	cH6–cH7	<1.0	1.2 ± 0.9
ω_8	terminal	aH7–aH8	9.6 ± 1.0	7.8 ± 0.6
	internal	bH7–bH8	<4.0	8.7 ± 0.1
	internal	cH7–cH8	<4.0	1.4 ± 0.7

Table 11: Experimental (ISPA) NOE Distances (Å) and Average Interproton Distances from MD and REMD Simulations for Fragments of *NmB* Trisialoside, during the First 70 ns and Last 30 ns of the Conventional MD Simulation^a

spins	NMR	MD	spins	NMR	MD
Preconformational Transition (The First 70 ns)					
bH3ax–cH8	2.6 ± 0.06	2.7 ± 1.0	bH6–bH8	2.3 ± 0.05	2.4 ± 0.2
bH3eq–cH8	3.5 ± 0.08	3.9 ± 0.5	aH3ax–bH8	2.8 ± 0.07	2.6 ± 1.0
bH3eq–cH7	3.9 ± 0.09	4.6 ± 0.4	aH3eq–bH8	3.4 ± 0.08	3.9 ± 0.5
bH6–bH7	2.2 ± 0.05	2.4 ± 0.1	bH9–cH7	3.1 ± 0.06	3.3 ± 1.8
Postconformational Transition (The Last 30 ns)					
bH3ax–cH8	2.6 ± 0.06	2.6 ± 1.0	bH6–bH8	2.3 ± 0.05	2.9 ± 0.3
bH3eq–cH8	3.5 ± 0.08	3.8 ± 0.4	aH3ax–bH8	2.8 ± 0.07	4.1 ± 0.2
bH3eq–cH7	3.9 ± 0.09	4.6 ± 0.4	aH3eq–bH8	3.4 ± 0.08	4.5 ± 0.1
bH6–bH7	2.2 ± 0.05	2.4 ± 0.1	bH9–cH7	3.1 ± 0.06	5.3 ± 1.5

^a The distances are averaged over the MD trajectories as $\langle 1/R^6 \rangle^{-1/6}$.

The interproton distances prior to and after the conformational transitions are presented in Table 11. It should be noted that the interproton distances were statistically indistinguishable in the b–c linkage for both segments of the trajectory. However, notable differences were observed in the a–b linkage. For example, the aH3ax–bH8 distance during the first 70 ns was 2.6 ± 1.0 Å, which compared well with the experimental value of 2.8 ± 0.07 Å, relative to the distance of 4.1 ± 0.2 Å computed during the last 30 ns. The bH6–bH8 and aH3eq–bH8 were also in better

Table 12: The Conformations (deg) of Representative Members of the Four Clusters in the Four-State Markov Model in the Methyl α -(2-8)-Disialoside

state	φ	ψ	ω	ω	populations from partition volumes (%) ^a
	(C1C2O8C8)	(C2O8C8C7)	8(O8C8C7O7)	7(O7C7C6O6)	
1 (red)	–97	71	73	65	16
2 (green)	–142	143	173	62	8
3 (blue)	–82	128	78	67	12
4 (yellow)	–167	106	74	69	64

^a Provides an estimate of what fraction of the data set belongs to a particular conformation. A total of 25 partition volumes were employed.

agreement with the experimental data during the first 70 ns of the simulation compared to the last 30 ns. Coupled with the J -values, these interproton distances indicate that during the last 30 ns of the conventional MD simulation experimentally nonrelevant conformations were sampled.

3.2. Conformational Clustering. In order to identify unique global conformational states and correlated internal rotations, the trajectories of **I** were subjected to a clustering analysis (40). Another key attribute of such an analysis is the accessibility of pertinent information about the interconversion pathways between conformational states, which cannot be easily discerned from traditional torsion angle scatter plots or from time dependent trajectory representations of dihedral angles. The conformer families were generated on the basis of inter-residue dihedral angles: φ , ψ , ω_8 and ω_7 .

A logarithmic plot of the fastest transition time scales τ_1 (relative to the time scale of the 25 dimensional model τ_{25}) for the hierarchy of Markov states obtained from the trajectory of **I** is shown in Figure 12A. The largest changes in time scale suggest a four-state model is adequate to describe the conformational space. The representative values of the glycosidic linkages of the four clusters present in the four-state model, and the abundance of each state, are presented in Table 12. Two states differing in more one than dihedral angle will effect transitions through correlated or concerted changes, while states with differences in a single angle undergo interconversions through isolated transitions. Transition probabilities were generally low in the case of strongly coupled or correlated transitions between two or more torsion angles, which may be ascribed to the overall additive barrier to interconversion. Examination of the four-

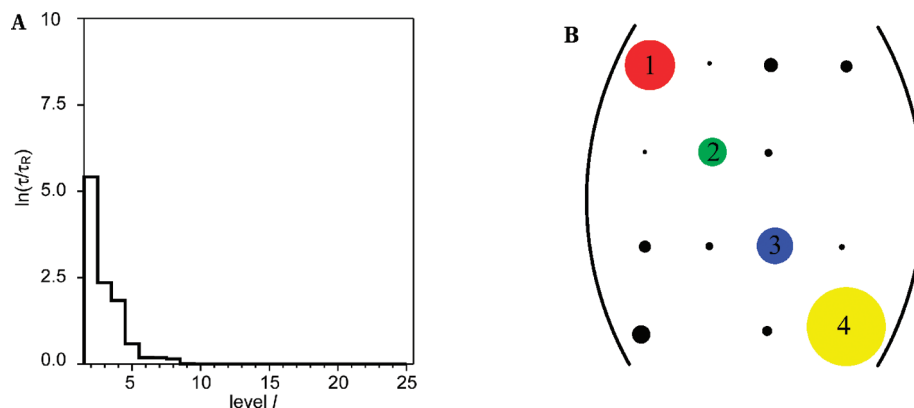


FIGURE 12: Plot of the relaxation time-scale, $\ln \tau/\tau_R$, between the various hierarchic Markov states identified from the MD-generated trajectories of the methyl α -(2,8)-disialoside, A. A graphical representation of the transition matrix between the Markov states when a four-state model is employed to describe the molecular conformational distribution of the dimer, B. The diameters of the diagonal peaks represent the populations of each state and cross-peaks denote populations of states and transition probabilities, respectively. The sizes of the cross-peaks are related to the additive energy barrier to interconversion.

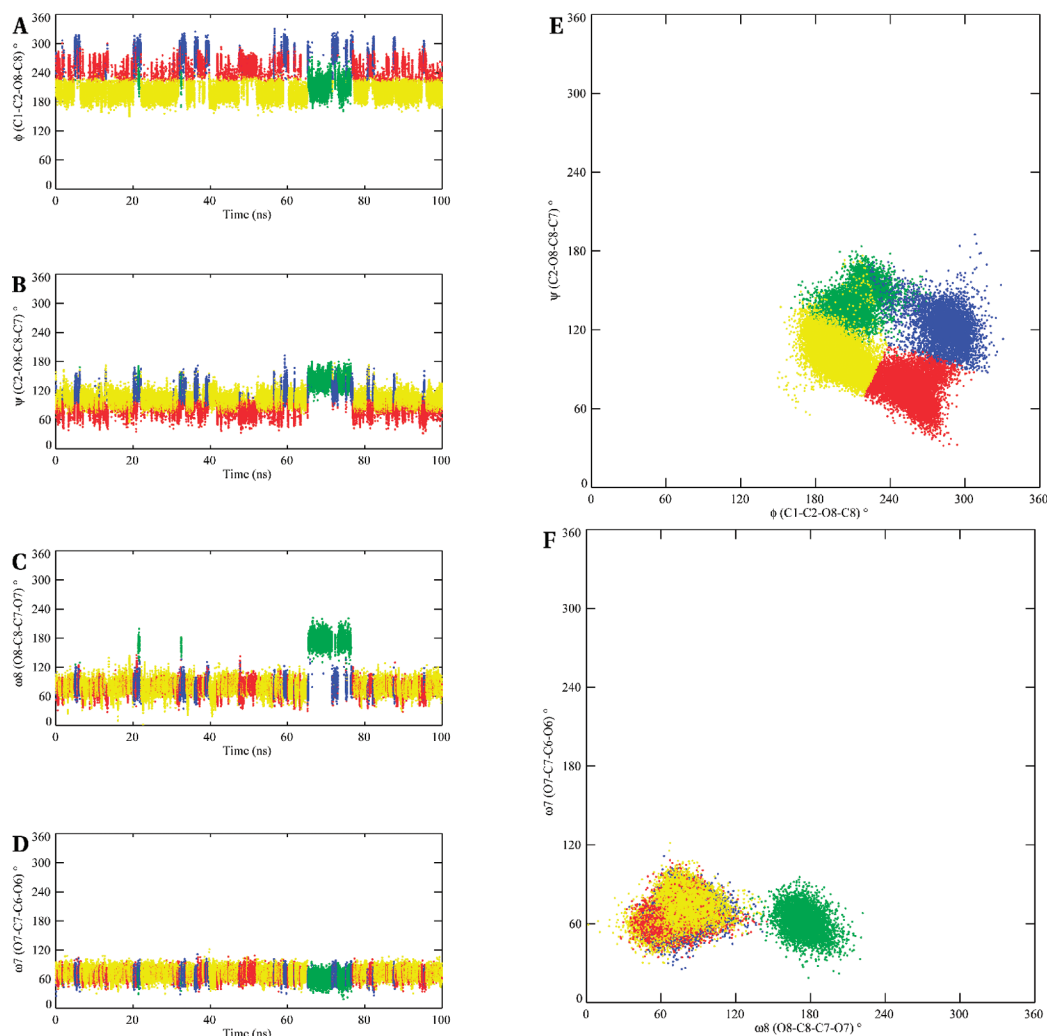


FIGURE 13: The conformational transition pathway of the four plausible clusters of methyl α -(2,8)-disialoside over the course of the MD simulation, left column. A, B, C and D represent the pathways of the φ , ψ , ω_8 and ω_7 angles, respectively. The scatter plots of the exocyclic dihedral angles, right column: E, (φ , ψ), and F, (ω_8 , ω_7). The figures employ the color-coding of Figure 12 to indicate the various conformations.

state transition matrix (Figure 12) indicates that state 1 (red) undergoes less frequent transitions to state 3 (blue) than to state 4 (yellow). The principal difference between states 1 and 3 lies in the ψ angles with values of 71° and 128° , respectively. States 1 and 4 differ in the (φ , ψ) domain with values of $(-97^\circ, 71^\circ)$ and $(-167^\circ, 106^\circ)$, respectively. Given that there are two concerted transitions in the 1–4 pathway, it might be expected that the 1–3 pathway would be preferred. However, the transitions between states 1 and 4 occur with higher probability. This discrepancy is explained in terms of the low barrier for the transitions between the *-gauche* and *anti* rotamers of the φ angle involved in the 1–4 route. State 2 (green) is characterized by an *anti* (173°) orientation of the ω_8 angle, and by an increase in the ψ angle (143°). This is an isolated state, as indicated by the extremely low transition frequency (Figure 12), which is accessed essentially only from state 3 (blue). The reason for its isolation arises in the need for simultaneous changes in ω_8 and ψ , to compensate for the electrostatic repulsions that would otherwise result from a single transition in ω_8 (Figure 8). The 1–2 pathway involves three concerted transitions (φ , ψ and ω_8) and would be disfavored, compared to two concerted changes (φ and ω_8) and (ψ and ω_8) in the 2–3 and 2–4 routes, respectively. The 2–3 pathway is more

probable compared to the 2–4 pathway, because the 2–3 route involves the φ angle oscillating frequently between the *-gauche* and *anti* rotamers. Pathways 3–4 and 1–4 both involve changes in the (φ , ψ) domain. However, the 3–4 route occurs with a lower probability. It must be the case that the 3–4 pathway is strongly competing with another pathway. Further examination indicated that state 3 also frequently interchanges with state 1, the (1–3) route involving only the ψ angle. In terms of populations, which are indicated by the number of partition volumes in Table 12, the isolated state 2 accounted for 8% of the conformer distribution. States 1 and 3 accounted for 16 and 12%, respectively, while state 4 was the dominant state, at 64%.

Plots of the time evolution of the glycosidic angles of the 4-state model during the simulation and scatter plots of the (φ , ψ) and (ω_8 , ω_7) dihedral angles are presented in Figure 10. During the first 65 ns and the last 20 ns of the simulation interconversions occurred predominantly between states 1 and 4, with much lower sampling of states 2 and 3, respectively. However, within the 65–80 ns interval, states 2 and 3 were preferentially populated. It should be noted that the major pathway from state 2 was via state 3. The scatter plots, Figure 13, indicate that the (φ , ψ)-space is

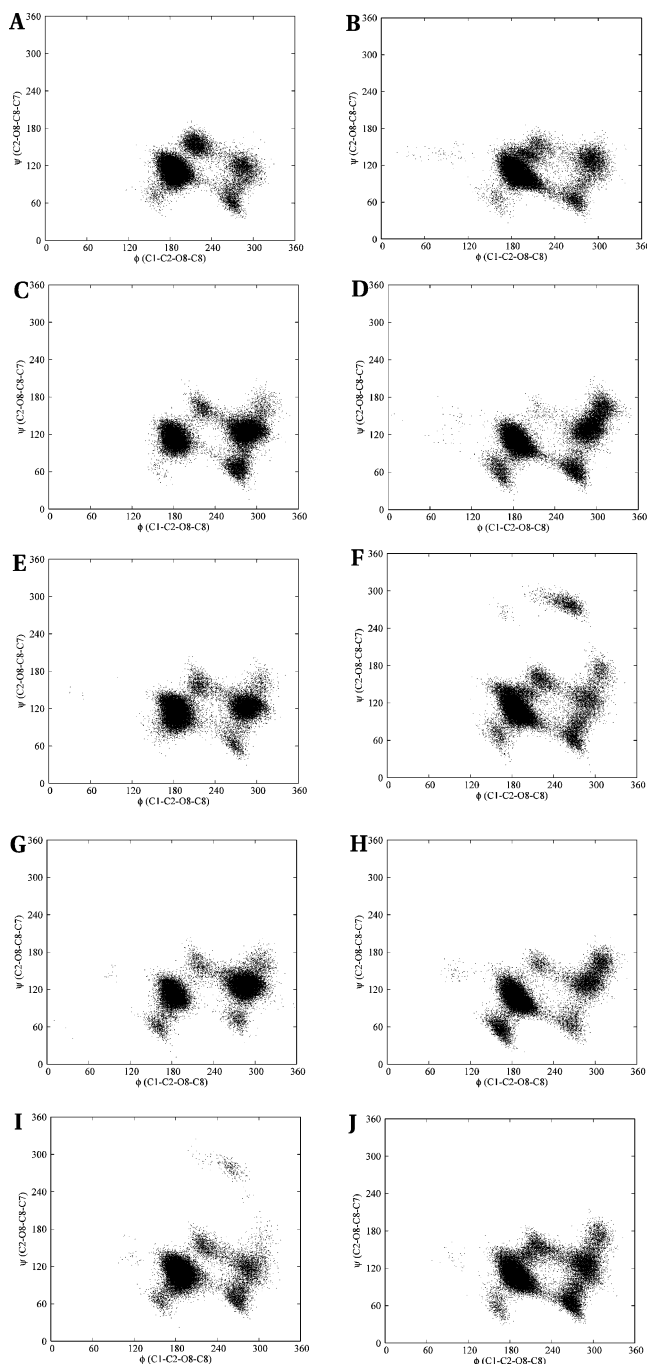


FIGURE 14: The (φ, ψ) conformational space of the native *N*-acetyl α -(2,8)-trisialoside and its analogues. The reducing and nonreducing linkages are on the left and right columns, respectively. *N*-acetyl (A, B), *N*-glycolyl (C, D), *N*-propionyl (E, F), *N*-propyl (G, H) and *N*-butanoyl (I, J).

clearly separated into four states for the 4-state model, while the (ω_8, ω_7) space showed an overlap of states 1, 3 and 4.

3.3. Effects of Chemical Group Modifications. The effects of chemical group modifications on the conformational properties of **II** were determined by comparing the (φ, ψ) and (ω_8, ω_7) scatter plots of the linkage dihedral angles with those of the *N*-Gc, *N*-Pr, *N*-Prop and *N*-But analogues. The multiple conformational distributions in the (φ, ψ) scatter plots indicate a significant degree of flexibility in the trisaccharide hapten and its analogues, Figure 14. For any given trisaccharide the φ and ψ angles for both the reducing and nonreducing terminal residues displayed similar distributions with the exception that the *N*-Pr and *N*-But analogues

occupied additional (φ, ψ) -spaces in the nonreducing and reducing linkage regions, respectively. Despite these differences in the number of conformations between the reducing and nonreducing linkage regions in each analogue, subsets of the phase-space were common to both linkage regions, albeit to varying extents. The (φ, ψ) -space of the native trimer, compared to all the analogues, exhibited similar conformational properties in terms of the number and conformation of the states, except for the *N*-Pr derivative, which displayed a significant population of a conformation with (φ, ψ) -values both between -120° and -60° . The (ω_8, ω_7) -space showed less flexibility in all cases, and considerable phase-space overlaps were seen in all the linkages of the trisaccharides, Figure 15. Among all the linkages the (ω_8, ω_7) -space showed more rigidity primarily due to the restricted flexibility of the ω_7 angle. However, in the *N*-Prop and *N*-But analogues the ω_7 angle populated isolated families with (ω_8, ω_7) values of about $(-300, 180)$ and $(60, 180)$, respectively. These additional conformational families were unexpected, given the preference for ω_7 to adopt a *+gauche* conformation in the native hapten.

In general the chemical group modifications resulted in similarities in the conformational families in the trisaccharides, although there were conformational states unique to some systems. To provide a structural interpretation for this observation, representative structures of conformational states identified via the Bayesian statistical method were generated. Plots of the relaxation time scales determined for each trajectory, Figure 16, indicated that the following number of states could adequately describe the conformational properties of each trisaccharide: **II** (10 states), *N*-Gc (16 states), *N*-Pr (16 states), *N*-Prop (14 states) and *N*-But (13 states). The representative conformations of the most populated states of the trisaccharide fragments are presented in Table 13. Based on dihedral angle matches, conformer family A of the *N*-Ac trimer will have the highest similarity with conformer family A of the *N*-Gc, *N*-Pr, *N*-Prop and *N*-But analogues, respectively. It is also evident that conformer family B of **II** and *N*-But trimers are close structural matches. However, these angle-matching comparisons may fail to identify certain global structural similarities. In particular, MD simulations have indicated that oligosaccharides may display similar overall topologies even though their internal linkages possess different dihedral angles (51), which are due to correlated transitions between the dihedral angles in the linkages. Therefore, the conformational similarities between the representative structures of the most populated clusters of the analogues and the native trisaccharide were also determined by rmsd comparisons. This approach revealed further structural matches that were not identified initially based on angle-similarities, Table 14 and Figure 17. For example, the A and J conformers of **II** and the *N*-Gc analogue, respectively, differ in the φ and ω_8 angles of the b–c linkage. Despite these differences, they share very similar overall structures, Figure 17. The relative abundance of the states, Table 14, was employed to determine the fraction of the states displaying similar overall topologies. In the case of **II** and the *N*-Gc trisialosides, 24% of the *N*-Gc conformers (A, C, J) exhibited similar backbone topologies as the A conformer of **II** (36%). Another 8% of the conformers of **II** (C) could be mapped to 12% of the *N*-Gc conformers (C), while 8% of the conformers of **II** (E) were

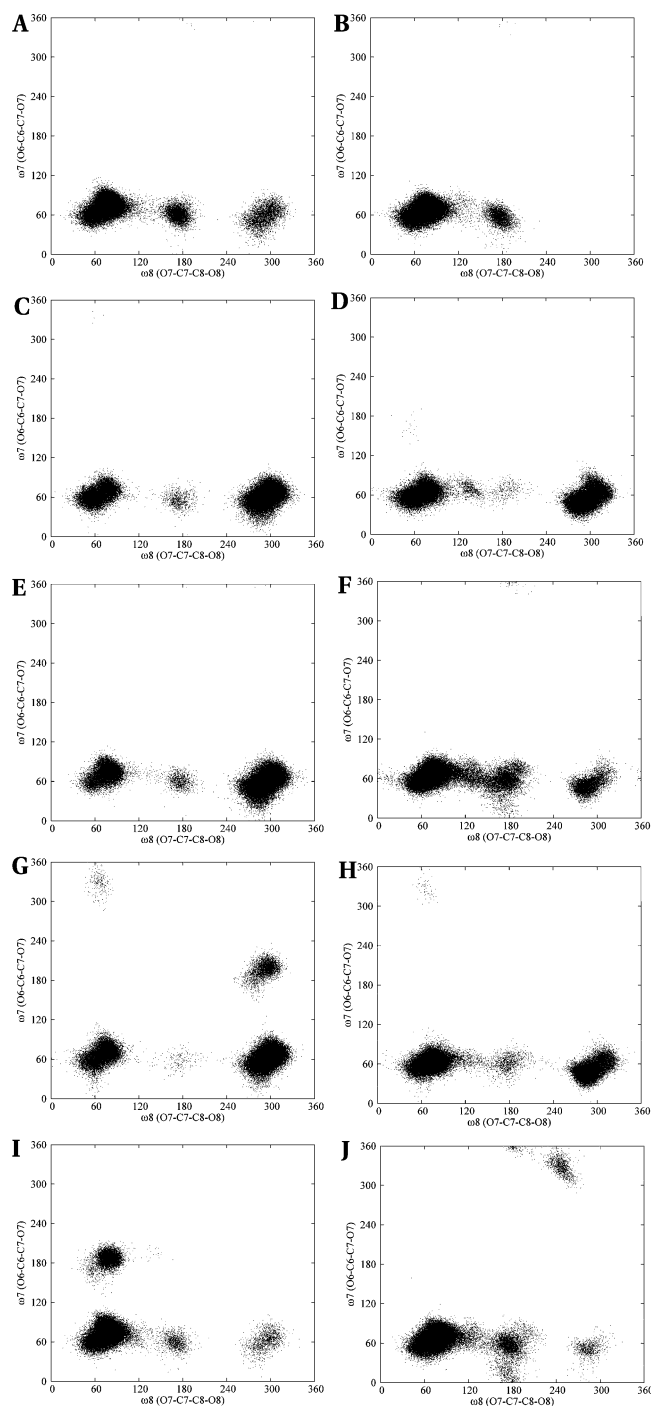


FIGURE 15: (ω_8 , ω_7) scatter plots of the conformational space of the native *N*-acetyl α -(2,8)-trisialoside and its analogues. The reducing and nonreducing linkages are in the left and right columns, respectively. *N*-Acetyl (A, B), *N*-glycolyl (C, D), *N*-propionyl (E, F), *N*-propyl (G, H) and *N*-butanoyl (I, J).

similar to 8% of the *N*-Glc conformers (I). Such comparisons were extended to all the other analogues (Table 14).

These topological similarities suggest that a common core conformational epitope is present in the *NmB* CPS and its analogues. These observations may be utilized to rationalize the results from efforts to enhance the immunogenicity of the *NmB* CPS by chemically modifying the *N*-acetyl moiety (11, 14, 18–22). In those studies, both IgM (20) and IgG (14, 18, 19) classes of antibodies, respectively, were utilized. Attempts to characterize the epitopes of the *N*-propionylated CPS suggested that the IgG mAbs recognized

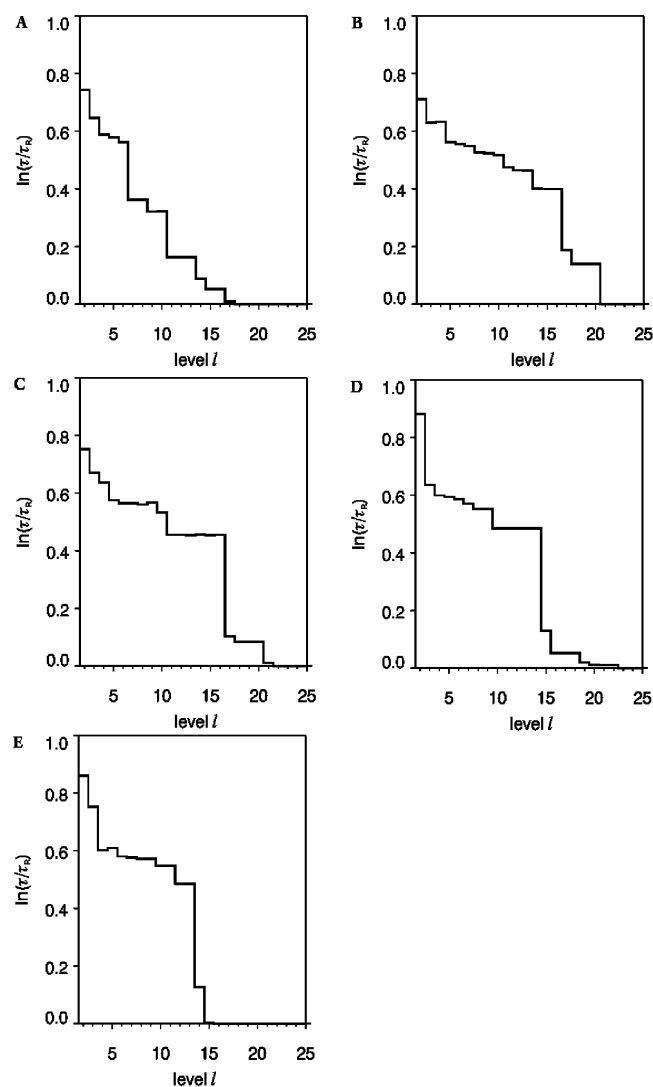


FIGURE 16: Plot of the relaxation time scale, $\ln(\tau/\tau_0)$, between the hierarchical Markov states identified from the REMD trajectories of II (A), *N*-Glc (B), *N*-Prop (C), *N*-Pr (D) and *N*-But (E).

both short and extended conformational epitopes (14), and a subset of the mAbs cross-reacted with human PSA and encapsulated *NmB* (14). In another study (20) IgM mAbs cross-reacted equally with the native *NmB* CPS and its derivatives, while different reactivities were noted among other species. In that study *NmB* CPS specific IgMs were bactericidal and safe when tested in mice. These studies indicate that low affinity IgMs and fully matured IgGs can show cross-reactivity between the native and modified *NmB* CPS. Regardless of the immunoglobulin class, the observed cross-reactivity or lack thereof may be rationalized in terms of interactions determined by the global (core) configuration of the polysaccharide, and those dependent on local interactions with pendant or exocyclic groups. Despite the observed similarities in backbone topologies, Figure 17, the exocyclic groups are displayed uniquely in each trisaccharide, Figure 18.

When cross-reactivity is observed among the altered CPS analogues, the elicited mAbs are recognizing a common topology, Figure 17, whose fine specificities do not likely depend on the *N*-acyl groups or on strong interactions with them. On the other hand, the absence of cross-reactivity arises either from significant variations in the overall 3D shape (unlikely based on the data presented here) or from incompatibilities with

Table 13: Linkage Torsion Angles of Representative Members of the Predominant Clusters Populated during the REMD Simulations of the Native Trimer and Its *N*-Acetyl Substituted Analogues (a–b–c–OMe)

conformer	a–b				b–c			
	φ (C1C2O8C8)	ψ (C2O8C8C7)	ω_8 (O8C8C7O7)	ω_7 (O7C7C6O6)	φ (C1C2O8C8)	ψ (C2O8C8C7)	ω_8 (O8C8C7O7)	ω_7 (O7C7C6O6)
<i>N</i> -acetyl								
A (36%)	–170	106	78	70	–174	115	78	75
B (8%)	–73	132	77	62	–174	112	76	74
C (12%)	–159	103	76	65	–82	96	76	67
D (12%)	–173	108	73	67	–131	147	–176	59
E (8%)	61	121	143	62	–164	128	–87	65
F (8%)	–157	141	179	54	–178	100	87	75
<i>N</i> -glycolyl								
A (8%)	–175	109	65	59	178	115	–67	60
C (8%)	–177	90	92	67	–115	81	36	60
G (8%)	–64	139	–49	59	–178	109	–73	61
H (16%)	–179	112	–67	54	–76	125	–62	63
I (8%)	–175	109	65	59	178	115	–67	60
J (8%)	–171	106	73	65	–72	123	–67	61
K (8%)	–73	137	–93	59	–94	131	–93	60
<i>N</i> -propionyl								
A (12%)	–173	106	76	69	–172	114	74	72
L (8%)	–174	133	153	53	–175	109	–67	60
M (8%)	–169	104	81	71	–76	120	–70	59
N (20%)	–172	108	73	65	–179	121	–71	56
O (8%)	–177	113	–68	54	–73	126	–65	62
<i>N</i> -propyl								
A (12%)	–169	101	78	65	–176	112	74	72
P (12%)	–160	108	–8	64	–81	127	96	62
Q (8%)	–170	101	77	61	–179	117	–67	61
R (16%)	–170	101	75	65	–75	125	–64	62
S (12%)	–177	113	–74	47	–73	128	–61	64
T (8%)	–176	113	–75	47	–71	132	–68	–164
<i>N</i> -butanoyl								
A (32%)	–172	113	75	66	–173	113	77	76
B (8%)	–74	118	77	64	–171	112	76	76
U (12%)	–85	112	131	46	–176	96	77	160
V (8%)	–94	71	69	62	–174	110	75	74
W (8%)	–174	72	60	57	–158	109	53	74

Table 14: Backbone^a RMSD (Å) Comparison between the Representative Conformations of the Native Trimer and Those of the Analogues, Populated during the REMD Simulations

	<i>N</i> -acetyl					
	A	B	C	D	E	F
<i>N</i> -glycolyl						
A	<i>0.79^b</i>	1.14	1.25	1.66	1.87	1.46
C	<i>1.00</i>	1.30	<i>0.69</i>	1.29	1.51	1.37
G	2.81	2.53	2.28	2.20	2.14	2.35
H	1.83	1.75	1.90	2.16	2.22	1.73
I	1.27	1.46	1.38	1.10	<i>1.26</i>	1.67
J	<i>1.39</i>	1.76	1.85	1.92	2.09	1.94
K	2.01	1.98	1.83	1.90	1.70	1.75
<i>N</i> -propionyl						
A	<i>0.72</i>	1.19	<i>1.21</i>	1.61	1.85	1.45
L	1.85	1.67	1.40	1.17	1.25	1.31
M	<i>1.50</i>	1.88	1.91	1.90	2.06	2.00
N	1.28	1.49	1.31	0.98	<i>1.18</i>	1.65
O	1.80	1.78	1.90	2.13	2.17	1.73
<i>N</i> -propyl						
A	<i>0.49</i>	1.29	1.61	1.91	2.03	1.58
P	1.86	1.78	1.32	1.22	1.67	2.15
Q	1.28	1.60	1.42	1.01	<i>1.17</i>	1.77
R	1.52	1.92	1.90	1.82	1.94	2.01
S	1.86	1.78	1.93	2.12	2.11	1.68
T	2.07	1.95	2.07	2.16	2.24	1.82
<i>N</i> -butanoyl						
A	<i>0.48</i>	1.00	1.18	1.60	1.83	1.40
B	1.17	<i>0.35</i>	1.23	1.67	2.02	1.23
U	1.22	1.10	1.63	1.80	2.04	1.37
V	0.87	0.92	1.39	1.75	2.13	1.59
W	0.84	1.40	1.55	1.86	2.08	1.84

^a Pyranose ring and glycosidic linkage atoms. ^b Italicized entries indicate the best structural matches (see also Figure 14).

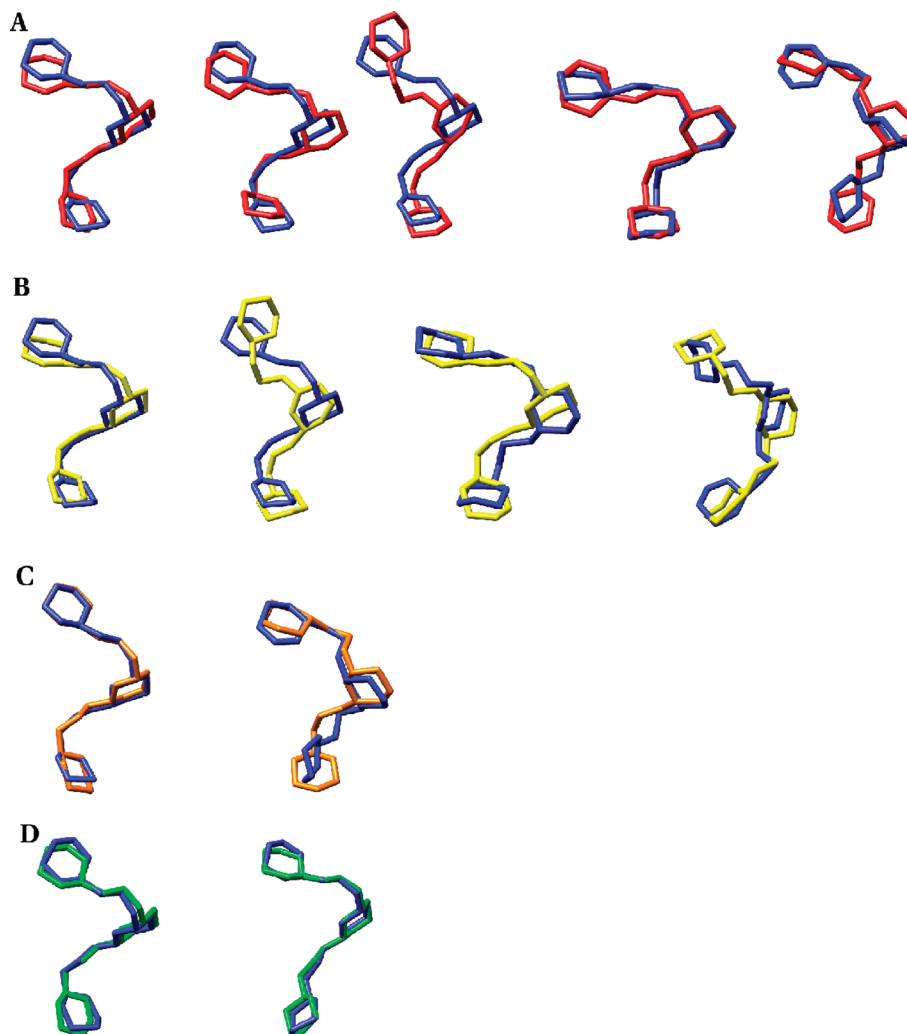


FIGURE 17: Backbone (ring and glycosidic linkage atoms) superposition of the representative members from the most populated clusters during the REMD simulations of the native and *N*-acetyl analogues of the methyl α -(2,8)-trisialoside. (A) *N*-Acetyl (blue) and *N*-glycolyl (red): A–A, A–C, A–J, C–C, E–I. (B) *N*-Acetyl (blue) and *N*-propionyl (yellow): A–A, A–M, C–A, E–N. (C) *N*-Acetyl (blue) and *N*-propyl (orange): A–A, E–Q. (D) *N*-Acetyl (blue) and *N*-butanoyl (green): A–A, B–B.

the presentation of pendant groups in the CPS. An antibody that has evolved to recognize the non-native *N*-acyl derivatives, such as mAb 13D9, may not be able to make optimal contacts when the native CPS is encountered, despite the presence of a common conformational backbone. This is particularly evident in the comparison between the native structure and the synthetic *N*-Pr or *N*-But analogues, Figure 18 and Figure 19. The smaller size of the *N*-acetyl group may prevent it from interacting effectively with a binding pocket deep enough to accommodate the larger *N*-Pr and *N*-But groups. This would explain the observation that although mAb 13D9 requires an extended epitope, as in the native CPS, it is unable to recognize the native CPS (23). It was reported previously that absence of cross-reactivity between the native CPS and the synthetic *N*-Pr and *N*-But polysaccharides could be due to conformational changes in the CPS induced by the modifications (20). In contrast, the MD simulations performed in this work suggest there is significant overlap in the conformational space of the native and chemically modified saccharides, observations that are in accord with experimental NMR data (11).

To illustrate the key structural properties pertinent to antibody recognition, decasaccharide helices were generated from structures selected from the conformational states identified during the REMD simulations of **II** and the *N*-But

analogue. The building blocks were selected to be consistent with the experimental NOE and *J*-value data. Conformational states whose φ -angles adopted the *anti* rotamer were selected because this rotamer preference led to the reproduction of the bH9–cH7 distance. States were also required to display the *+gauche* rotamers of ω_8 and ω_7 , consistent with a strong NOE between H8–H6, and low magnitude 3J -values between H8–H7 and H7–H6. Based on these criteria, state A was selected for both **II** and its *N*-But analogue. As expected, the NOE distances computed from these states are consistent with the NMR data, Table 15. The helical parameters, *n* (the number of residues per turn) and *h* (vertical rise per residue in Å), were calculated based on the method described by Miyazawa et al. (52). Vectors consisting of the glycosidic oxygen atoms (O8) were employed to describe virtual bonds in each decasaccharide. Using these vector distances, angles and dihedral angles for the central residues (residues 4 to 7) were computed for each decasaccharide and employed to determine *n* and *h*. The (*n*, *h*) values were (2.2, 4.5 Å) and (2.3, 5.2 Å) for the A state of **II** and its *N*-But analogue, respectively. It should be noted that, in solution, these polymers are dynamic and interconvert between states, and these idealized helices are not representative of the overall structure. Nonetheless, the helices generated here, Figure 19,

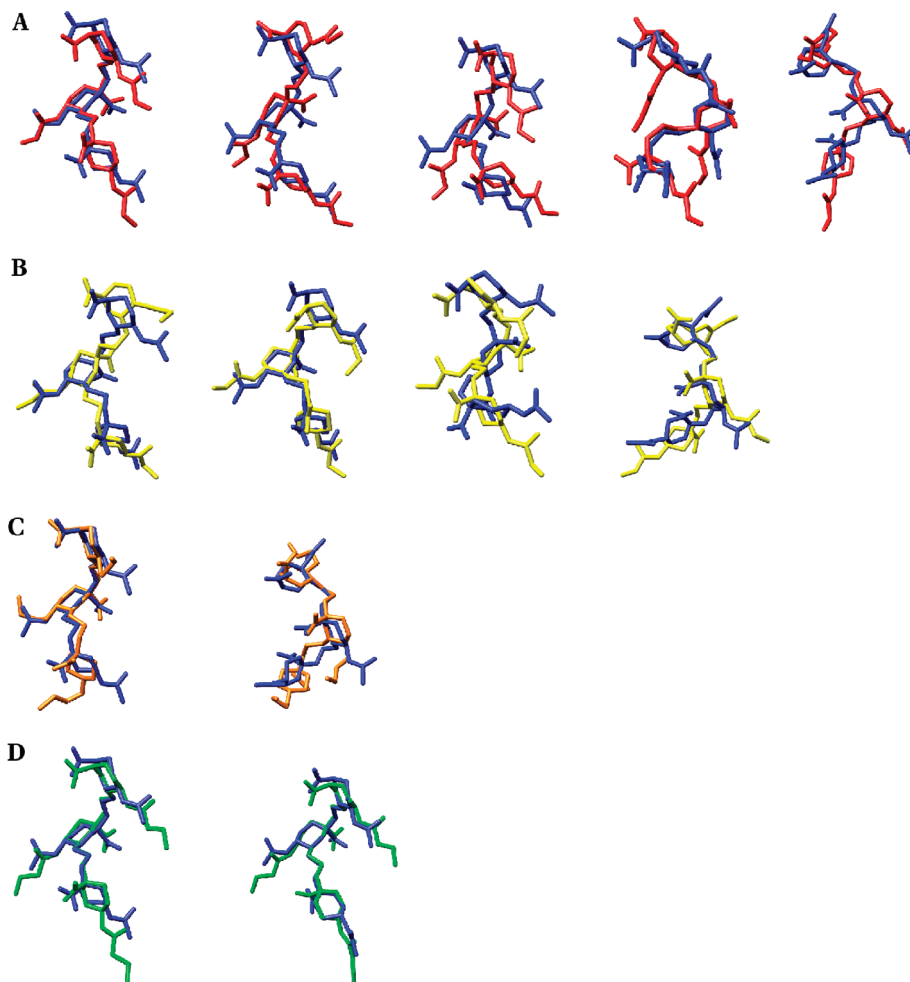


FIGURE 18: The display of key exocyclic groups relative to the backbone (ring and glycosidic linkage atoms) conformations of the representative cluster members of the native methyl α -(2,8)-trisialoside and its analogues. (A) *N*-Acetyl (blue) and *N*-glycolyl (red): A–A, A–C, A–J, C–C, E–I. (B) *N*-Acetyl (blue) and *N*-propionyl (yellow): A–A, A–M, C–A, E–N. (C) *N*-Acetyl (blue) and *N*-propyl (orange): A–A, E–Q. (D) *N*-Acetyl (blue) and *N*-butanoyl (green): A–A, B–B.

are representative of the local epitopes that are presented to antibodies. The outward projection of the *N*-acyl groups from the helical axis is evident, suggesting that these groups may be crucial for antibody recognition. In these helices the *N*-acetyl and *N*-butanoyl groups constitute 33% and 46% of their respective surfaces. Consequently, an antibody whose specificity involves the *N*-butanoyl moiety would be expected to have much lower affinity for the *N*-acetyl analogue. In each case, the carboxylate groups are also presented on the helical surfaces, although in the *N*-But analogue these groups are markedly less accessible.

Two salient points emerge from all these considerations: cross-reactivity arises between the *N*-acyl analogues when overall backbone conformation is conserved and when the immune response has not matured to recognize the acyl moiety as an immunodominant epitope. Therefore, these simulations suggest that alternative chemical modifications may be sought, subject to the constraints that the integrity of the *N*-acetyl moiety, and the backbone core conformations, are not compromised. These modifications could involve the introduction of small hydrophobic groups at positions that are hydrophilic in the native structure, and thus less immunogenic. To the best of our knowledge, no such derivatizations have been performed with the *NmB* CPS. However, within the accuracy limitations of the force field and MD simulations, it is now feasible to

predict the effect of chemical modifications on the CPS structure and to identify the structural similarities between native and synthetic immunogens.

4. CONCLUSION

Because of the absence of a protective vaccine against *NmB*, and the postulated importance of a conformational epitope in mediating the antigenicity and bactericidal activity related to this CPS, considerable attention has been given to characterizing its 3D shape and dynamics (9, 11, 18, 25, 26, 46). In addition, the weak immunogenicities of both the protein-conjugated and nonconjugated forms of the *NmB* CPS have resulted in alternative vaccine design attempts that typically involve chemical group modifications of the acyl groups (11, 14, 18, 21, 22) or the carboxylate group (11).

As an initial step in quantifying the antigenicities of native and chemically modified *NmB* CPS, conformational properties of di- and trisaccharide haptens have been determined by MD simulations and confirmed by NMR spectroscopy. That the MD simulations identified experimentally relevant conformations was demonstrated by the good agreement between the experimental and theoretical NMR observables. In addition, a Bayesian statistical analysis (40) was employed to identify conformational states from the trajectories. To

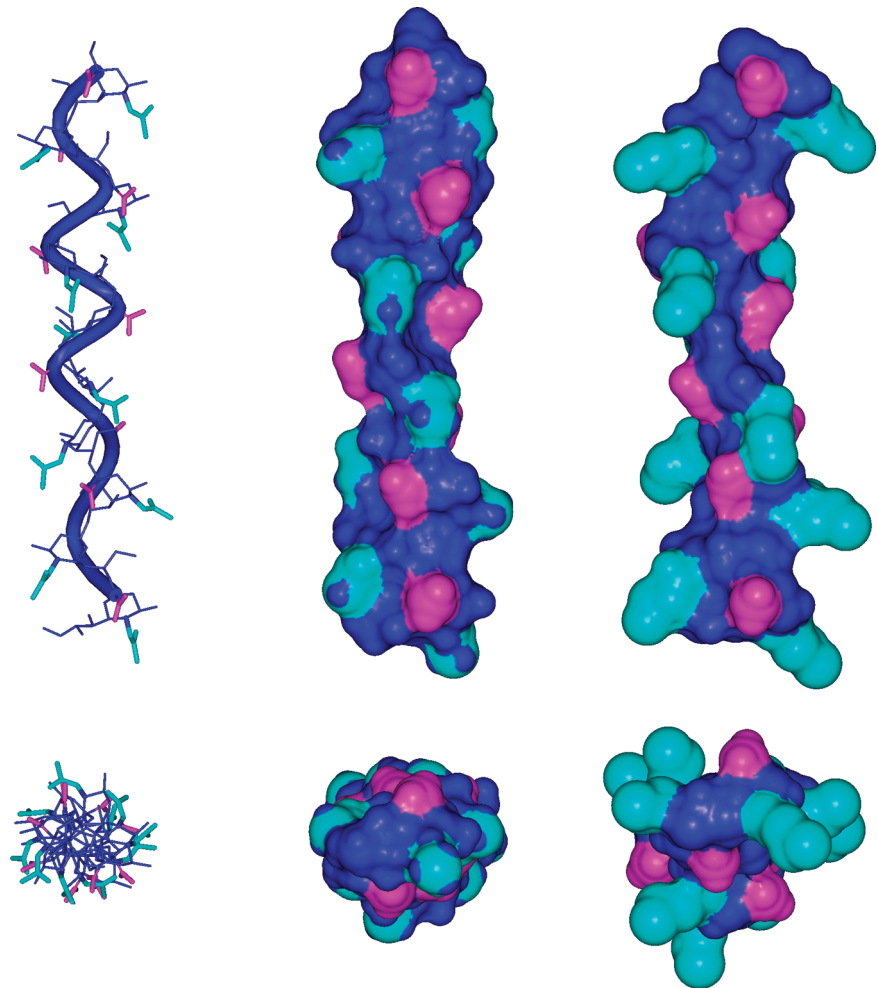


FIGURE 19: Longitudinal and axial views of deca-saccharide helical structures generated by propagating representative conformational states derived from the REMD simulations of the native trisaccharide (left and middle panels) and the *N*-butanoyl analogue (right panel). The helical repeat is eight residues in each helix, shown to overlap in the axial views. Cyan, *N*-acyl groups; pink, carboxyl groups; blue, all other atoms.

Experimental (ISPA) NOE and MD-Derived Distances (Å)							
spins	NMR (native)	<i>N</i> -acetyl	<i>N</i> -butanoyl	spins	NMR (native)	<i>N</i> -acetyl	<i>N</i> -butanoyl
bH3ax–cH8	2.6 ± 0.06	2.2 ± 0.2	2.3 ± 0.2	bH6–bH8	2.3 ± 0.05	2.4 ± 0.2	2.4 ± 0.2
bH3eq–cH8	3.5 ± 0.08	3.5 ± 0.2	3.6 ± 0.2	aH3ax–bH8	2.8 ± 0.07	2.2 ± 0.2	2.3 ± 0.2
bH3eq–cH7	3.9 ± 0.09	5.1 ± 0.2	5.1 ± 0.2	aH3eq–bH8	3.4 ± 0.08	3.6 ± 0.2	3.6 ± 0.2
bH6–bH7	2.2 ± 0.05	2.4 ± 0.1	2.4 ± 0.1	bH9–cH7	3.1 ± 0.07	2.8 ± 0.4	2.9 ± 0.4

Experimental and MD-Derived ³ <i>J</i> -Values					
angle	linkage	spins	NMR (native)	<i>N</i> -acetyl	<i>N</i> -butanoyl
ω_7	internal	bH6–bH7	<1.0	1.0 ± 0.7	1.1 ± 0.9
	internal	cH6–cH7	<1.0	1.5 ± 1.0	1.6 ± 1.0
ω_8	internal	bH7–bH8	<4.0	1.2 ± 0.7	1.2 ± 0.7
	internal	cH7–cH8	<4.0	1.2 ± 0.8	1.1 ± 0.6

^a REMD simulations were performed at 8 temperatures for a total of 160 ns. The distances are averaged over the MD trajectories as $\langle 1/R^6 \rangle^{(-1/6)}$.

the best of our knowledge this is the first time such an analysis has been applied to investigate carbohydrate conformational properties. With the preliminary agreement demonstrated here between the MD data and the experimental data for the native haptens, the study was extended to shed light on the effects of chemical group modifications on epitope size and fine structure. These theoretical simulations present a unique opportunity to generate atomic level insight into the conformational space of native and synthetic CPS

analogues. It is also suggested that efforts employed in the quest for more immunogenic carbohydrate vaccines that can mimic *NmB* should consider chemical group modifications that do not perturb the *N*-acetyl moieties or the global conformation. MD simulations provide a much needed tool for the rational design of such synthetic vaccines, and provide a solid basis for the interpretation of the diverse and complex data from immunological studies.

ACKNOWLEDGMENT

R.J.W. thanks H. J. Jennings (NRC, Canada) and J. B. Robbins (NIH) for useful discussions.

SUPPORTING INFORMATION AVAILABLE

Pyranose ring vicinal proton–proton coupling constants of methyl α -(2,8)-di/trisialoside. This material is available free of charge via the Internet at <http://pubs.acs.org>.

REFERENCES

1. Ferreiros, C. M., Gomez, J. A., and Criado, M. T. (1998) Prevention of meningococcal disease: present and future. *Rev. Med. Microbiol.* 9, 29–37.
2. Tikhomirov, E., Santamaria, M., and Esteves, K. (1997) Meningococcal disease: public health burden and control. *World Health Stat. Q.* 50, 170–177.
3. Tsang, R. S. W., and Zollinger, W. D. (2005) Serological specificities of murine hybridoma monoclonal antibodies against *Neisseria meningitidis* serogroups B, C, Y and W135 and evaluation of their usefulness as serogrouping reagents by indirect whole-cell enzyme-linked immunosorbent assay. *Clin. Diagn. Lab. Immunol.* 12, 152–156.
4. Bardotti, A., Averani, G., Berti, F., Berti, S., Galli, C., Giannini, S., Fabbri, B., Proietti, D., Ravenscroft, N., and Ricci, S. (2005) Size determination of bacterial capsular oligosaccharides used to prepare conjugate vaccines against *Neisseria meningitidis* groups Y and W135. *Vaccine* 23, 1887–1899.
5. Bhattacharjee, A. K., Jennings, H. J., Kenny, C. P., Martin, A., and Smith, I. C. P. (1976) Structural determination of polysaccharide antigens of *Neisseria meningitidis* serogroup Y, serogroup W-135, and serogroup BO. *Can. J. Biochem.* 54, 1–8.
6. Bhattacharjee, A. K., Jennings, H. J., Kenny, C. P., Martin, A., and Smith, I. C. P. (1975) Structural determination of sialic acid polysaccharide antigens of *Neisseria meningitidis* serogroup B and serogroup C with C-13 nuclear magnetic resonance. *J. Biol. Chem.* 250, 1926–1932.
7. Bundle, D. R., Smith, I. C. P., and Jennings, H. J. (1974) Determination of the structure and conformation of bacterial polysaccharides by carbon 13 nuclear magnetic resonance. Studies on group-specific antigens of *Neisseria meningitidis* serogroups A and X. *J. Biol. Chem.* 249, 2275–2281.
8. Kabat, E. A., Liao, J., Osseman, E. f., Gamian, A., Michon, F., and Jennings, H. J. (1988) The epitope associated with the binding of the capsular polysaccharide of the group B meningococcus and of *Escherichia coli* K1 to a human monoclonal macroglobulin, IGM^{NOV}. *J. Exp. Med.* 168, 699–711.
9. Brisson, J. R., Baumann, H., Imbert, A., Perez, S., and Jennings, H. J. (1992) Helical epitope of the group B meningococcal α -(2-8)-linked sialic acid polysaccharide. *Biochemistry* 31, 4996–5004.
10. Jennings, H. J., Roy, R., and Michon, F. (1985) Determinant specificities of the group B and group C polysaccharides of *Neisseria meningitidis*. *J. Immunol.* 134, 2651–2657.
11. Baumann, H., Brisson, J. R., Michon, F., Pon, R., and Jennings, H. J. (1993) Comparison of the conformation of the epitope of α -(2-8) polysialic acid with its reduced and *N*-acyl derivatives. *Biochemistry* 32, 4007–4013.
12. Weintraub, A. (2003) Immunology of bacterial polysaccharide antigens. *Carbohydr. Res.* 338, 2539–2547.
13. Finne, J., Leinonen, M., and Makela, P. H. (1983) Antigenic similarities between brain components and bacteria causing meningitis. Implications for vaccine development and pathogenesis. *Lancet* 2, 355–357.
14. Garonoff, D. M., Bartoloni, A., Ricci, S., Gallo, E., Rosa, D., Ravenscroft, N., Guarnieri, V., Seid, R. C., Shan, A., Usinger, W. R., Tan, S., McHugh, Y. E., and Moe, G. R. (1998) Bactericidal monoclonal antibodies that define unique meningococcal B polysaccharide epitopes that do not cross-react with human polysialic acid. *J. Immunol.* 160, 5028–5036.
15. Devi, S. J. N., Robbins, J. B., and Schneerson, R. (1991) Antibodies to poly[(2-8)- α -N-acetylneuraminic acid] and poly[(2-9)- α -N-acetylneuraminic acid] are elicited by immunization of mice with *Escherichia coli* K92 conjugates: Potential vaccines for groups B and C meningococci and *E. coli* K1. *Proc. Natl. Acad. Sci. U.S.A.* 88, 7175–7179.
16. Lively, M. R., Gilbert, A. S., and Moreno, C. (1981) Sialic acid polysaccharide antigens of *Neisseria meningitidis* and *Escherichia coli*. Esterification between adjacent residues. *Carbohydr. Res.* 94, 193–203.
17. Azurmendi, H. F., Vionnet, J., Wrightson, L., Trinh, L. B., Shiloach, J., and Freedberg, D. I. (2007) Extracellular structure of polysialic acid explored by on cell solution NMR. *Proc. Natl. Acad. Sci. U.S.A.* 104, 11557–11561.
18. Evans, S. V., Sigurskjold, B. W., Jennings, H. J., Brisson, J. R., To, R., Tse, W. C., Altman, E., Frosch, M., Weisgerber, C., Kratzin, H. D., Klebert, S., Vaesen, M., Bittersuermann, D., Rose, D. R., Young, N. M., and Bundle, D. R. (1995) Evidence for the extended helical nature of polysaccharide epitopes. The 2.8Å resolution structure and thermodynamics of ligand binding of an antigen binding fragment specific for α -(2-8)-polysialic acid. *Biochemistry* 34, 6737–6744.
19. Jennings, H. J., Roy, R., and Gamian, A. (1986) Induction of meningococcal group B polysaccharide-specific IgG antibodies in mice by using an *N*-propionylated B polysaccharide-tetanus toxoid conjugate vaccine. *J. Immunol.* 137, 1708–1713.
20. Lively, M. R., and Esdaile, J. (1991) Specificity of the immune response to the group B polysaccharide of *Neisseria meningitidis*. *Immunology* 74, 490–496.
21. Pon, R. A., Khieu, N. H., Yang, Q. L., Brisson, J. R., and Jennings, H. J. (2002) Serological and conformational properties of *E. coli* K92 capsular polysaccharide and its *N*-propionylated derivative both illustrate that induced antibody does not recognize extended epitopes of polysialic acid: Implications for a comprehensive conjugate vaccine against groups B and C *N. meningitidis*. *Can. J. Chem.* 80, 1055–1063.
22. Pon, R. A., Lussier, M., Yang, Q. L., and Jennings, H. J. (1997) *N*-propionylated group B meningococcal polysaccharide mimics a unique bactericidal capsular epitope in group B *Neisseria meningitidis*. *J. Exp. Med.* 185, 1929–1938.
23. Pon, R. A., Biggs, N. J., and Jennings, H. J. (2007) Polysialic acid bioengineering of neuronal cells by *N*-acyl sialic acid precursor treatment. *Glycobiology* 17, 249–260.
24. Lively, M. R., Lindon, J. C., Williams, J. M., and Moreno, C. (1985) Structural and conformational features of the *Escherichia coli* K92 capsular polysaccharide. *Carbohydr. Res.* 143, 191–205.
25. Henderson, T. J., Venable, R. M., and Egan, W. (2003) Conformational flexibility of the group B meningococcal polysaccharide in solution. *J. Am. Chem. Soc.* 125, 2930–2939.
26. Venable, R. M., Bizik, F., Henderson, T. J., and Egan, W. (1997) Molecular dynamics simulations of an α -(2-8)-linked sialic acid tetramer in vacuum and solvent. *J. Mol. Struct. (THEOCHEM)* 395, 375–388.
27. Cumming, D. A., and Carver, J. P. (1987) Virtual and Solution Conformations of Oligosaccharides. *Biochemistry* 26, 6664–6676.
28. Gonzalez-Outeirino, J., Kadirvelraj, R., and Woods, R. J. (2005) Structural elucidation of type III group B Streptococcus capsular polysaccharide using molecular dynamics simulations: the role of sialic acid. *Carbohydr. Res.* 340, 1007–1018.
29. Vasudevan, S. V., and Balaji, P. V. (2002) Molecular dynamics of alpha2-8 linked disialoside: Conformational analysis and implications for binding to proteins. *Biopolymers* 63, 168–180.
30. Kirschner, K. N., and Woods, R. J. (2001) Solvent interactions determine carbohydrate conformation. *Proc. Natl. Acad. Sci. U.S.A.* 98, 10541–10545.
31. Gonzalez-Outeirino, J., Kirschner, K. N., Thobhani, S., and Woods, R. J. (2006) Reconciling solvent effects on rotamer populations in carbohydrates - A joint MD and NMR analysis. *Can. J. Chem.* 84, 569–579.
32. Kirschner, K. N., Yongye, A. B., Gonzalez-Outeirino, J., Tschampel, S. M., Daniels, C. R., Foley, B. L., and Woods, R. J. (2008) GLYCAM06: A Generalizable Biomolecular Force Field. *Carbohydrates. J. Comput. Chem.* 29, 622–655.
33. Yongye, A. B., Foley, B. L., and Woods, R. J. (2008) On achieving experimental accuracy from molecular dynamics simulations of flexible molecules: aqueous glycerol. *J. Phys. Chem. A* 112, 2634–2639.
34. Case, D. A., Darden, T. A., Cheatham, T. E., III, Simmerling, C. L., Wang, J., Duke, R. E., Lou, R., Merz, K. M., Wang, B., Pearlman, D. A., Crowley, M., Brozell, S., Tsui, V., Gohlke, H., Mongan, J., Hornak, V., Cui, G., Beroza, P., Schafmeister, P., Caldwell, J. W., Ross, W. S., and Kollman, P. A. (2004) AMBER 8, University of California, San Francisco, CA.
35. Jorgensen, W. L., and Swenson, C. J. (1985) Optimised intermolecular potential functions for amides and peptides. Structure and properties of liquid amides. *J. Am. Chem. Soc.* 107, 569–578.

36. Darden, T., York, D., and Pederson, L. (1993) Particle mesh Ewald: A $N \log(N)$ method for Ewald sums in large systems. *J. Chem. Phys.* 98, 10089–10092.
37. van Gunsteren, W. F., and Berendsen, H. J. C. (1977) Algorithms for macromolecular dynamics and constraint dynamics. *Mol. Phys.* 34, 1311–1327.
38. Okur, A., Wickstrom, L., Layten, M., Geney, R., Song, K., Hornak, V., and Simmerling, C. (2006) Improved efficiency of replica exchange simulations through use of a hybrid explicit/implicit solvation model. *J. Chem. Theory Comput.* 2, 420–433.
39. Rathore, N., Chopra, M. J., and dePablo, J. (2005) Optimal allocation of replicas in parallel tempering simulations. *J. Chem. Phys.* 122, 024111.
40. Schultheis, V., Hirschberger, T., Carstens, H., and Tavan, P. (2005) Extracting Markov models of peptide conformational dynamics from simulation data. *J. Chem. Theory Comput.* 1, 515–526.
41. Kloppenburg, M., and Tavan, P. (1997) Deterministic annealing for density estimation by multivariate normal mixtures. *Phys. Rev. E* 55, R2089–R2092.
42. James, T. L., and Leu, H. E. (1999) *CORMA*, University of California, San Francisco.
43. Thrippleton, M. J., and Keeler, J. (2003) Elimination of zero-quantum interference in two-dimensional NMR spectra. *Angew. Chem., Int. Ed.* 42, 3938–3941.
44. Delaglio, F., Grzesiek, S., Vuister, G. W., Zhu, G., Pfeifer, J., and Bax, A. (1995) Nmrpipe - A Multidimensional Spectral Processing System Based On Unix Pipes. *J. Biomol. NMR* 6, 277–293.
45. NMRView, NMRView Software; One Moon, Scientific, Inc.
46. Michon, F., Brisson, J. R., and Jennings, H. J. (1987) Conformational differences between linear α -(2-8)-linked homosialooligosaccharides and the epitope of the group B meningococcal polysaccharide. *Biochemistry* 26, 8399–8405.
47. Haasnoot, C. A. G., Deleeuw, F., and Altona, C. (1981) Relationship between Proton-Proton NMR Coupling-Constants and Substituent Electronegativities. I. An empirical generalization of the Karplus Equation. *Tetrahedron* 36, 2783–2792.
48. Kooijman, H., Kroonbatenburg, L. M. J., Kroon, J., Breg, J. N., and Deboer, J. L. (1990) Structure of α -D-N-acetyl-1-O-methylneuraminic acid methyl ester. *Acta Crystallogr., Sect. C: Cryst. Struct. Commun.* 46, 407–410.
49. Poppe, L., Stuike-Prill, S., Meyer, B., and van Halbeek, H. (1992) The solution conformation of sialyl- α -(2,6)-lactose studied by modern NMR techniques and Monte Carlo simulation. *J. Biomol. NMR* 2, 109–136.
50. Breg, J., Kroon-Batenburg, M. J. L., Strecker, G., Montreuil, J., and Vliegthart, J. F. G. (1989) Conformational analysis of the sialyl- α -(2–3/6) N-acetylactosamine structural element occurring in glycoproteins, by two-dimensional nOe ^1H -NMR spectroscopy in combination with energy calculations by hard-sphere exoanomeric and molecular mechanics force-field with hydrogen-bonding potential. *Eur. J. Biochem.* 178, 727–739.
51. Woods, R. J., Pathiaseril, A., Wormald, M. R., Edge, C. J., and Dwek, R. A. (1998) The high degree of internal flexibility observed for an oligomannose oligosaccharide does not alter the overall topology of the molecule. *Eur. J. Biochem.* 158, 372–386.
52. Sugeta, H., and Miyazawa, T. (1967) General method for calculating helical parameters of polymer chains from bond lengths, bond angles and internal rotation angles. *Biopolymers* 5, 673–679.

BI800431C