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MM3 MODELING OF ALDOPENTOSE PYRANOSE RINGS

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ABSTRACT

MM3 (version 1992, $\epsilon = 3.0$) was used to study the ring conformations of D-xylopyranose, D-lyxopyranose and D-arabinopyranose. The energy surfaces exhibit low-energy regions corresponding to chair and skew forms with high-energy barriers between these regions corresponding to envelope and half-chair forms. The lowest energy conformer is 4C_1 for α - and β -xylopyranose and α - and β -lyxopyranose, and the lowest energy conformer is 1C_4 for α - and β -arabinopyranose. Only α -lyxopyranose exhibits a secondary low-energy region (1C_4) within 1 kcal/mol of its global minimum. Overall, the results are in good agreement with NMR and crystallographic results. For many of these molecules, skew conformations are found with relatively low energies (2.5 to 4 kcal/mol above lowest energy chair form). The 2S_0 and 1C_4 conformers of crystalline benzoyl derivatives of xylopyranose are in secondary low-energy regions on the β -xylopyranose surface, within 3.8 kcal/mol of the global 4C_1 minimum.

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INTRODUCTION

Previously, we modeled the pyranosyl ring shapes of the D-aldohexoses with the molecular mechanics program MM3.^[1] In general, MM3 predicted the shapes of these cyclic structures well. All of the β -anomers favored the 4C_1 conformation, as did most of the α -anomers. For the hexoses experimentally found in multiple forms, e.g. α -idopyranose and α -altropyranose, the model predicted multiple forms. However, the conformer distribution predicted for these molecules was not always in exact agreement with the distributions predicted from NMR results, signifying the effects of modeling error, including neglected solvation and entropic effects. In addition, a comparison of the predicted energies for the anomers indicated that the equatorially configured β -structures were systematically overpredicted.

A better test of a potential energy function for predicting carbohydrate ring forms would be a series of pyranosyl structures known to be more variable in conformational preference. The aldopyranosyl pentoses, which lack the bulky hydroxymethyl group, represent such a series. By NMR,^[2,3] α - and β -xylopyranose and β -lyxopyranose exist predominately in the 4C_1 form; α - and β -arabinopyranose exist predominately in the 1C_4 form; and α -lyxopyranose exists as a mixture of the two chair forms. Ribopyranose, which has already been studied with MM3,^[4] is also found as a mixture of chair forms. This series appears to be a good test of the ability of potential functions to reproduce ring conformations.

In addition to the goal of validating molecular models, there is interest in understanding how saccharides interact with proteins and how these interactions facilitate enzyme kinetics. In part, this stems from the need to improve enzyme stability or to alter reaction conditions. Xylanase is of commercial interest because it reduces the need for chlorine bleaching in paper pulping.^[5,6] A number of xylanase structures with xylan fragments or related derivatives bound to the active site have now been solved by diffraction crystallography, and these structures suggest that bound xylose rings can have non-chair conformations during hydrolysis.^[7-9] An understanding of the conformational preferences of xylopyranose may be important for understanding the activity and specificity of these enzymes.

METHODS

The computational methods used in this work have been described in previous publications and will only be outlined here.^[1,4] To be consistent with those calculations, the 1992 version of MM3^[10-12] was used with an elevated dielectric constant ($\epsilon = 3.0$). The elevated dielectric constant was used to reduce the strength of intramolecular hydrogen bonding and better mimic carbohydrates in condensed-phase systems.^[13] Planar ring structures were initially generated with PC-Model (Serena Software, Bloomington, IN). These structures were then used to create puckered rings by moving three alternating ring atoms and their connected substituents perpendicular to the initial ring plane. This was accomplished by orienting the planar ring into the xy -plane and adjusting the z -coordinates of the atoms to be shifted. The atoms were moved over a range of -0.8 \AA to $+0.8 \text{ \AA}$ in 0.1 \AA increments, which was sufficient to cover all likely ring shapes. For each ring, all three staggered gauche conformations of the four second-

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dary hydroxyl groups were also varied. Combined with the different combinations of ring shape, this yielded a total of 397,872 ($17^3 \times 3^4$) starting structures for each monosaccharide. Ring pucker was maintained during the energy minimization by fixing the z -coordinates of the six ring atoms. All other degrees of freedom were allowed to relax. These optimizations were conducted with the block-diagonal Newton–Raphson method (MM3 Option 1) and the default optimization criteria. The system of Cremer and Pople (q, ϕ, θ) was used to quantify ring shape.^[14]

From the full set of optimized structures, a Pickett–Strauss *plate carrée* representation (ϕ, θ) of the Cremer and Pople puckering sphere was generated.^[15] To construct this two-dimensional representation, the data was sorted into $10^\circ \times 10^\circ$ regions of ϕ and θ , regardless of the value of q or orientation of the individual exocyclic groups. The lowest energy conformer from each region was selected, and the resulting grid of points was plotted using SURFER (Golden Software, Golden, CO). Positions of the characteristic conformers (chairs, boats, skews, etc.) within this two-dimensional space are shown in Figure 1.

The Pickett–Strauss surface distorts some aspects of the puckering sphere. Most affected are the two poles (i.e. the chair conformers), which are transformed into the top and bottom y -axes. In effect, the points representing the poles are stretched into lines corresponding to the two horizontal boundaries. Because the system used to generate the puckered rings evenly samples the puckering sphere, the transformation results in a low density of data points near the two chair regions. This results in some distortion of the contour lines near these boundaries.

To find local minima, the local low-energy regions were identified from the Pickett–Strauss surfaces. Molecules with these ring shapes were built with all 81 (3^4) sets of staggered exocyclic orientations. All of these structures were freely optimized to determine the lowest-energy form. If any of the structures from one local region optimized to a different area of the puckering space, the region was not considered to have a local minimum. The Karplus equations of Haasnoot et al.^[16] were used to calculate proton–proton coupling constants for these structures.

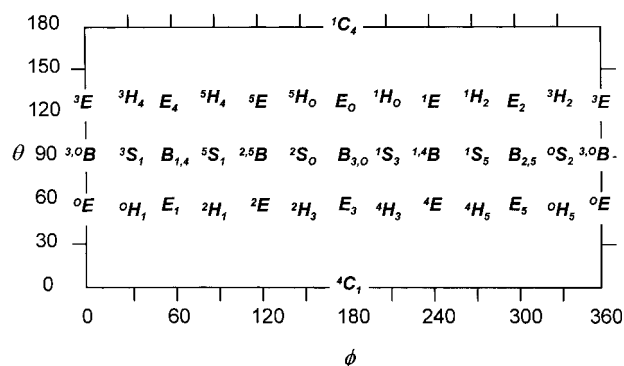


Figure 1. Location of the characteristic chair, boat, skew, half-chair, and envelope forms of the aldopyranosyl monosaccharides on the Pickett–Strauss energy surface (ϕ, θ).

The Cambridge Crystal Structure Database^[17] was searched to identify related crystal structures. If there were multiple entries within the database, only the most recent determination was included in the analysis, and neutron diffraction determinations were used in preference to x-ray determinations. Structures for compounds containing L-sugars were included after converting the puckering parameters ($\phi_D = 180^\circ - \phi_L$, $\theta_D = 180^\circ - \theta_L$) to account for the change in absolute configuration. The Protein Data Bank (18) was searched for structures with bound xylose rings.

RESULTS AND DISCUSSION

All three pyranosyl energy surfaces (Figures 2–4) have low-energy regions corresponding to chair and skew forms. The half-chair and envelope forms are relatively high energy and form barriers between the low-energy regions. The MM3 minima for these monosaccharides are reported in Table 1, and selected low-energy structures are shown in Figure 5.

For both α - and β -xylopyranose, the lowest-energy ring conformer is 4C_1 . The second-lowest energy conformer for α -xylopyranose is 1C_4 and for β -xylopyranose is

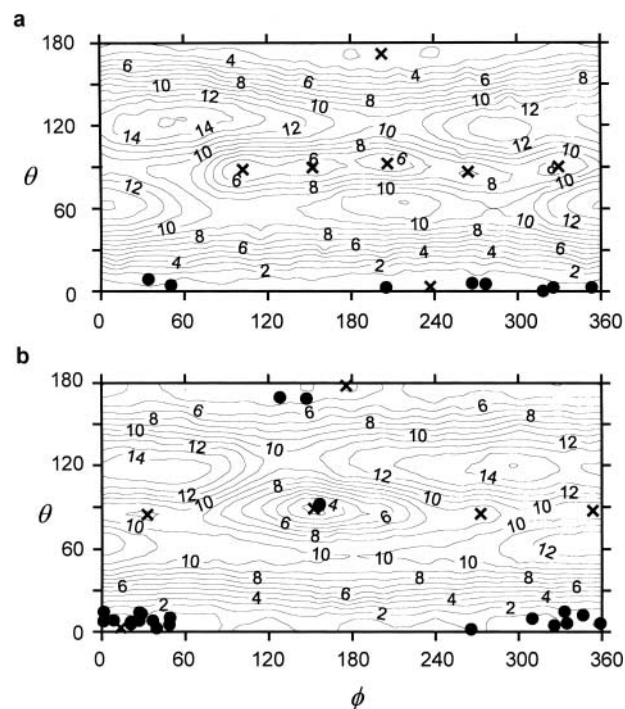


Figure 2. Ring puckering energy surfaces for D-xylopyranose: (a) α -xylopyranose and (b) β -xylopyranose. Local minima are labeled with crosses (x). Ring shapes of crystal structures are labeled with circles (o).

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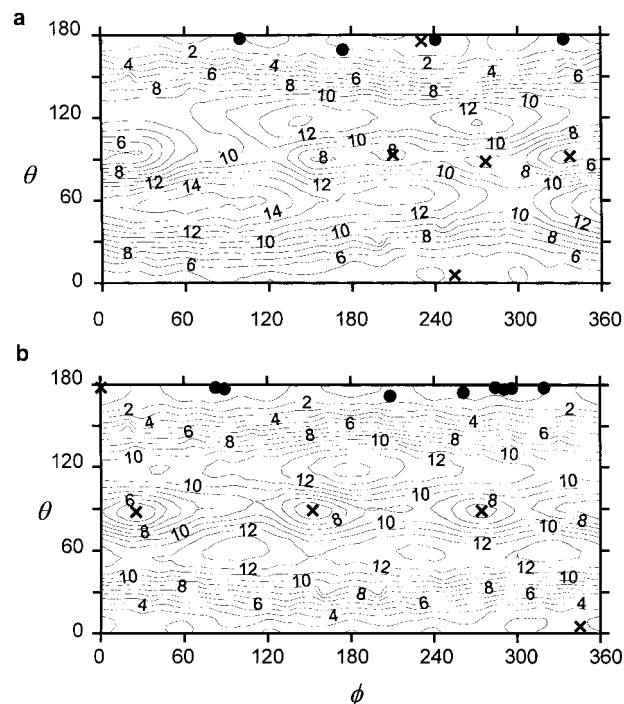


Figure 3. Ring puckering energy surfaces for D-arabinopyranose: (a) α -arabinopyranose and (b) β -arabinopyranose. Local minima are labeled with crosses (x). Ring shapes of crystal structures are labeled with circles (o).

2S_0 . These forms are 1.7 and 2.6 kcal higher in energy, respectively. The 1C_4 form of β -xylopyranose is the third lowest energy form at 3.8 kcal/mol. The energy differences between the two chair forms are much smaller for xylopyranose than for glucopyranose,^[1] demonstrating the large energy penalty for having the bulky hydroxymethyl side group in an axial orientation. Nevertheless, because the 4C_1 conformation dominates the distribution, the calculated coupling constants for the 4C_1 form and the conformational average are similar. Both sets of coupling constants are in good agreement with NMR results (Table 1), with a root-mean-square (rms) deviation between the calculated and experimental coupling constants of 0.6 Hz for α -xylopyranose and 0.4 Hz for β -xylopyranose.

Numerous single crystal structures exist for molecules that contain xylopyranosyl rings.^[19-42] Most of these structures have 4C_1 forms (Figure 2). Benzoyl derivatives, however, are exceptions.^[28,29] For these molecules 2S_0 and 1C_4 ring shapes are found, apparently because of stabilization induced by intramolecular stacking of the benzoyl rings in these conformations. Both ring shapes correspond to secondary minima on the β -xylopyranosyl MM3 energy surface (Figure 2b). The experimental observation of these alternative conformers for xylose derivatives supports the relatively low energies calculated by MM3 for these regions.

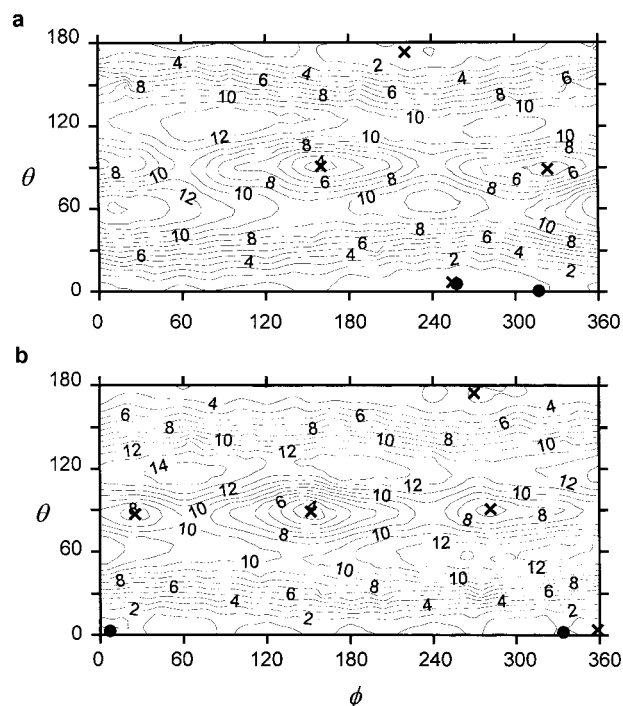


Figure 4. Ring puckering energy surfaces for D-lyxopyranose: (a) α -lyxopyranose and (b) β -lyxopyranose. Local minima are labeled with crosses (\times). Ring shapes of crystal structures are labeled with circles (\bullet).

In addition to single crystal structures, a number of enzymes have been crystallized with bound xylosyl moieties.^[7-9] Although bound subunits are generally less accurately determined in protein diffraction studies than in small-molecule diffraction studies, non-chair xylose ring shapes have been reported in these structures by different research groups.^[8,9] In these studies, the xylose ring in the -1 subsite of the family 11 xylanases was observed in a ${}^{2.5}B$ conformation. This conformation appears to facilitate the formation of an oxo-carbenium ion transition intermediate, which is key to the proposed double displacement catalytic mechanism that accounts for the retention of the β -xylose configuration during hydrolysis. The observed ${}^{2.5}B$ boat can be obtained from the relatively low-energy 2S_O form by the downward movement of the anomeric carbon as it forms the covalent xylose-glutamic acid bridge of the intermediate. Small downward shifts of the O-5 and C-2 ring atoms and the C-2 and C-3 hydroxyl groups are required for this conformational change, but no movement of the C-4 oxygen and its covalently bound oligosaccharide moiety is needed. In terms of puckering parameters, the ϕ angle is reduced from $\sim 150^\circ$ in the 2S_O form to $\sim 120^\circ$ in the ${}^{2.5}B$ form (Figure 1), and the MM3 energy change for this conversion is only 2-3 kcal/mol (Figure 2b). Sidhu et al.^[9] have proposed an enzyme-mediated ring conversion pathway of ${}^4C_1 \rightarrow {}^2H_3 \rightarrow {}^2S_O \rightarrow {}^{2.5}B$ for the formation of this intermediate.

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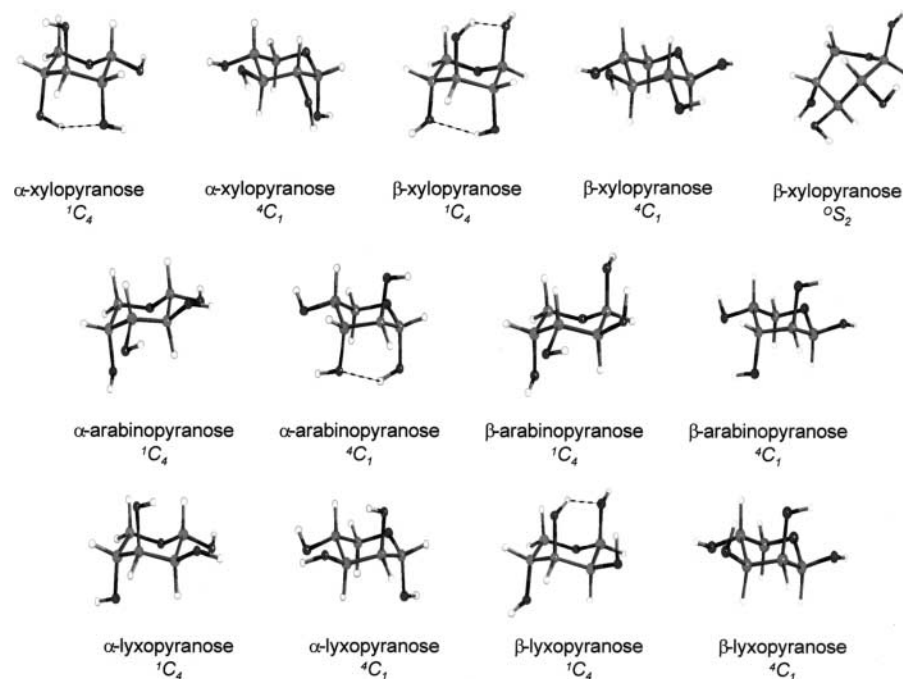


Figure 5. Selected low-energy chair and skew conformers for D-xylopyranose, D-arabinopyranose, and D-lyxopyranose. Intramolecular hydrogen bonds are shown as dashed lines ($H \cdots O$ distance > 2.0 Å).

However, the relatively low energy of the 2S_0 form (2.6 kcal/mol) may indicate that this conformer exists in solution in a concentration sufficient to allow for the direct binding of this form to the enzyme. In this instance, the enzyme would not need to induce the conformational change through the 2H_3 form, which has a much higher relative energy.

Both anomers of arabinopyranose have 1C_4 lowest energy conformations and 4C_1 second lowest-energy conformations (Figure 3). For the β -anomer, the 4C_1 form, which has the C-1 hydroxyl group in an equatorial position, is only 1.45 kcal/mol above the global minimum region. Consequently, a small amount ($< 10\%$) of this form may exist in solution. The NMR coupling constants calculated for the 1C_4 chair and the conformational equilibrium are in good agreement with experimental values (Table 1). The rms deviation between the experimental and calculated coupling constants for the α - and β -anomers are 0.6 and 0.4 Hz, respectively. In addition, all reported arabinopyranosyl crystal structures^[43–52] have 1C_4 conformations (Figure 3).

The α -D-lyxopyranose surface has a 4C_1 global low-energy region and a 1C_4 secondary low-energy region (Figure 4A). The energy difference between these two forms is 0.9 kcal/mol (Table 1), indicating that both of the chairs are likely to exist in equilibrium. Except for J_{23} , which has a value of ~ 3 Hz for either chair, the reported coupling constants are between the values for the two chair forms, which also indicates

Table 1. MM3 Energies, Puckering Parameters, and Proton-Proton Coupling Constants for the MM3 Local Minima of Xylopyranose, Lyxopyranose and Arabinopyranose

Conformer	Puckering Parameters			MM3 Energies, kcal/mol		x_i^a	Coupling Constants, Hz				
	q , Å	θ , deg	ϕ , deg	Steric	Relative		J_{12}	J_{23}	J_{34}	J_{45d}	J_{45u}
<i>α-xylopyranose</i>											
4C_1	0.561	3.42	236.87	8.935	0.000	0.949	3.5	9.0	8.6	10.6	5.9
1C_4	0.558	172.01	202.47	10.668	1.733	0.050	1.2	3.4	2.9	2.2	1.1
5S_1 - ^{25}B	0.723	88.06	102.07	13.807	4.872	0.000	2.5	7.0	0.3	2.8	0.7
1S_3	0.780	92.30	206.08	13.945	5.009	0.000	3.3	5.4	9.1	7.2	9.4
2S_0	0.762	89.52	152.04	14.003	5.068	0.000	5.2	9.9	5.3	1.2	4.2
1S_5	0.743	86.45	264.18	15.865	6.930	0.000	1.2	0.2	6.3	10.6	4.4
0S_2	0.729	90.14	329.47	16.224	7.289	0.000	3.5	4.8	0.8	8.1	9.1
Conformational ave. Exp. ^b				9.025			3.4	8.8	8.3	10.1	5.7
							3.7	9.8	9.1	10.2	5.1
<i>β-xylopyranose</i>											
4C_1	0.591	3.01	13.52	7.939	0.000	0.987	7.2	9.0	8.5	10.6	6.0
2S_0	0.765	88.86	152.14	10.557	2.618	0.012	4.5	9.9	5.6	1.2	4.4
1C_4	0.519	178.07	175.91	11.748	3.809	0.002	2.5	3.1	3.1	2.2	1.2
5S_1	0.713	85.19	272.57	16.390	8.451	0.000	2.9	0.4	5.2	10.6	4.7
3S_1	0.738	84.67	32.46	16.768	8.829	0.000	5.5	0.3	3.8	1.3	5.1
0S_2 - ^{30}B	0.737	87.38	353.47	17.148	9.210	0.000	1.3	3.7	2.5	5.2	9.3
Conformational ave. Exp. ^b				7.975			7.1	9.0	8.5	10.5	6.0
							7.8	9.2	9.0	10.5	5.6
<i>α-D-lyxopyranose</i>											
4C_1	0.550	6.83	254.12	9.429	0.000	0.825	2.3	3.1	8.8	5.8	10.6
1C_4	0.568	173.09	220.88	10.353	0.924	0.172	7.3	3.1	3.2	1.1	2.2
2S_0	0.756	90.76	160.11	12.893	3.464	0.002	0.8	2.1	6.3	5.3	1.4
0S_2	0.750	88.80	323.65	13.670	4.240	0.001	5.0	2.3	0.4	8.5	9.2



that both chairs co-exist in solution. Based on the MM3 energies of the local minima, the model predicts an 83%/17% distribution of the 4C_1 and 1C_4 forms, which appears to underpredict the 1C_4 form. As a result, the rms deviation between the calculated and experimental coupling constants is relatively large (1.3 Hz). A least squares best-fit model of the two chairs to the experimental coupling constants improves the fit (0.7 Hz deviation) and yields a 62%/38% ratio of the 4C_1 and 1C_4 forms. Crystalline acetyl and methyl derivatives of α -D-lyxopyranose^[53,54] are both found in 4C_1 conformations. MM3 calculations for β -D-lyxopyranose suggest that the 4C_1 form is the most likely ring shape. The calculated NMR coupling constants are in reasonable agreement with experimental results (0.7 Hz rms deviation), and the crystal structures for β -L-lyxopyranose^[55] and methyl-2,3,4-triacetyl- β -D-lyxopyranoside^[56] are also found in the 4C_1 form. The second low-energy region for β -D-lyxopyranose is the 1C_4 , which is 2.6 kcal/mol above the global minimum.

Anomeric ratios can be estimated from the average energies of the conformer populations for each monosaccharide. For xylopyranose, the distribution is calculated to be 14% α -anomer and 86% β -anomer. Compared with the experimental ratio in D_2O (37% α -anomer/63% β -anomer),^[3] the β -anomeric form is overpredicted. For lyxopyranose, the calculated distribution is 48% α -anomer and 52% β -anomer, and again the β -form is overpredicted compared to the distribution in D_2O (71% α -anomer/29% β -anomer).^[3] A similar difference was previously found to occur for the aldohexoses (1). For arabinopyranose, the calculated distribution is 82% α -anomer and 18% β -anomer. In this case, however, the β -anomer is underpredicted in comparison to NMR results in D_2O (63% α -anomer/37% β -anomer).^[3]

The systematic difference in the predicted anomeric ratios appears to indicate a small error in the MM3 potential function. For most of the hexoses, and for xylose and lyxose, MM3 energies overpredict the β -pyranosyl form. For these molecules, both anomers are found predominately in the 4C_1 form by calculation. For arabinose, MM3 underpredicts the β -pyranosyl form, and in this case both anomers exist predominately in the 1C_4 form by calculation. For idopyranose and altropyranose, the MM3 calculated anomer distributions appear to be very close to experimental values. These molecules model as a conformational equilibrium that includes significant amounts of both chair forms, which appears to dampen the effect. All of the overpredicted forms, β -anomers for structures in 4C_1 and α -anomers for structures in 1C_4 , have equatorial orientations of the glycosidic C-1—O bond and *trans* orientations of this bond relative to the C-5—O-5 bond. The underpredicted forms, in contrast, all have axial orientations of the glycosidic hydroxyl group and *gauche* orientations relative to the C-5—O-5 bond. Because the torsional functions associated with the anomeric sequences (O—C—O—H and O—C—O—C) have been especially difficult to model, this functionality is likely responsible for the calculated differences. The analysis, however, is complicated because MM3 also overpredicts the axial form of 2-hydroxytetrahydropyran relative to experiment (1). Recent ab initio results highlight these effects, and some improvement in the modeling of the anomeric region for disaccharides has been achieved by a hybrid QM/MM approach that in principle corrects for errors in the molecular mechanics modeling of anomeric sequences.^[57] In any case, the magnitude of the energy discrepancy in the MM3 calculation is small and is on the order of magnitude (<1 kcal/mol) of the uncertainty of the quantum mechanical calculations.



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There has been considerable debate regarding the importance of solvent and entropic effects on the conformation of small carbohydrates. Inclusion of these “additional” influences is often used to compensate for discrepancies between experimental data and the underlying computational method. With MM3, monosaccharide rings are modeled reasonably well without needing to incorporate other effects. The only accommodation to environmental forces is an increase in the bulk dielectric constant. The principal effect of this is to reduce the strength of intramolecular hydrogen bonds, which may reduce entropy differences that arise because of preferred intramolecular hydrogen bond orientations. In addition, although some effort was made to limit the influence of intermolecular effects in parameterizing MM3, the parameters are derived from a wide variety of experimental and computational data and some external influences may have crept into the model. Consequently, it cannot be absolutely concluded that intermolecular effects and entropy differences are small for the various conformers of these structures.

CONCLUSIONS

MM3 (version 1992, $\epsilon = 3.0$) was used to study the ring puckering of the two anomeric forms of D-xylopyranose, D-lyxopyranose, and D-arabinopyranose. The modeling indicates that α - and β -xylopyranose and β -lyxopyranose exist predominately in the 4C_1 form, that α - and β -arabinopyranose exists predominately in the 1C_4 form, and that α -lyxopyranose exists as an equilibrium between the two chair forms. The predicted conformers are in good agreement with those expected from NMR coupling constants and crystallographic data. For the pentoses, secondary conformations are lower in energy than they were for the hexoses, demonstrating the influence of the C-6 hydroxymethyl group on hexose conformation. A systematic error in the estimation of anomer ratios appears to be related to difficulties in parameterizing some torsion sequences.

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