

Modeling of negative Poisson's ratio (auxetic) crystalline cellulose $\ensuremath{\mathsf{I}\beta}$

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ORIGINAL PAPER



Modeling of negative Poisson's ratio (auxetic) crystalline cellulose $I_{\mbox{\scriptsize B}}$

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Abstract Energy minimizations for unstretched and stretched cellulose models using an all-atom empirical force field (molecular mechanics) have been performed to investigate the mechanism for auxetic (negative Poisson's ratio) response in crystalline cellulose I_{β} from kraft cooked Norway spruce. An initial investigation to identify an appropriate force field led to a study of the structure and elastic constants from models employing the CVFF force field. Negative values of on-axis Poisson's ratios v_{31} and v_{13} in the x_1-x_3 plane containing the chain direction (x_3) were realized in energy minimizations employing a stress perpendicular to the hydrogen-bonded cellobiose sheets to simulate swelling in this direction due to the kraft cooking process. Energy minimizations of structural evolution due to stretching along the x_3

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Materials and Engineering Research Institute, Sheffield Hallam University, Howard Street, Sheffield S1 1WB, UK e-mail: A.Alderson@shu.ac.uk chain direction of the 'swollen' (kraft cooked) model identified chain rotation about the chain axis combined with inextensible secondary bonds as the most likely mechanism for auxetic response.

Keywords Cellulose · Auxetic · Negative Poisson's ratio · Polymers

Introduction

Interest in determining the structure and properties of crystalline cellulose is due to the wide-ranging applications for cellulose and also for the potential to develop synthetic derivatives. Cellulose is a linear polymer formed by β -1,4-linked D-glucopyranose residues with high crystallinity, and has two main crystalline forms (cellulose I and cellulose II). Cellulose I has two crystalline allomorphs, designated cellulose I_{α} and I_{β} (Atalla and Van der Hart 1984; Van der Hart and Atalla 1984).

Interestingly, experimental measurements of a negative value of Poisson's ratio have been reported in crystalline cellulose II (Nakamura et al. 2004) and also in kraft cooked Norway spruce (Peura et al. 2006), which comprises crystalline cellulose I_{β} . A negative value of Poisson's ratio corresponds to auxetic behavior (Evans et al. 1991) and is characterized by a transverse expansion upon axial stretching of the material. Negative Poisson's ratios have also been

reported in a Raman spectroscopy study of the deformation of fibrous networks of bacterial cellulose and microcrystalline cellulose filaments (Tanpichai et al. 2012) and in a variety of paper and paperboard grades (fibrous materials consisting of self-binding cellulose fibers) (Stenberg and Fellers 2002; Verma et al. 2014). The origin of the auxetic nature of the fibrous networks has been attributed to the fiber network structure containing bent fibers forming reentrant structures, facilitated by strong hydrogen bonds at the junctions where fibers overlap (Tanpichai et al. 2012; Verma et al. 2014). In the case of the kraft cooked Norway spruce study, the negative Poisson's ratio in crystalline cellulose I_{β} was identified as v_{31} , corresponding to an increase in the inter-planar separation (along the x_1 direction) of planar sheets (in the x_2-x_3 plane) of parallel cellobiose chains for a tensile load applied along the cellobiose chain axis (x_3) direction. Peura et al. (2006) suggested the auxetic effect may be attributed to non-alignment of microfibrils leading to a component of the stretching force acting perpendicular to the hydrogen-bonded sheets, and shearing effects between individual cells and cell wall layers.

Auxetic materials are attracting interest for their apparently anomalous behavior, and also because the ability to tailor materials to display negative Poisson's ratio response can open up routes to extreme values of other properties not available for non-auxetic materials (Evans 1990; Greaves et al. 2011). Examples of enhanced properties as a consequence of the auxetic effect include fracture toughness (Lakes 1987), double (synclastic) curvature under pure bending (Lakes 1987; Evans 1991), indentation resistance (Alderson et al. 1994), shear rigidity, (Choi and Lakes 1992) and ultrasonic (Alderson et al. 1997) and vibration (Howell et al. 1991; Scarpa et al. 2005) damping.

Auxetic metals, polymers, composites and ceramics are known, from the macroscale to the nanoscale, and include natural as well as man-made materials (Greaves et al. 2011; Alderson and Alderson 2007; Evans and Alderson 2000). In terms of auxetic polymers, macroporous thermoplastic (Lakes 1987) and thermoset (Friis et al. 1988) foams have been developed, as have microporous polymers in the form of tape or ribbon (Caddock and Evans 1989), monolithic cylinders and plaques (Alderson and Evans 1992), and also in monofilament (Alderson et al. 2002) and film (Ravirala et al. 2005) form. Locally auxetic behavior has also been reported at the sub-micron scale in elastomeric polypropylene (Franke and Magerle 2011). At the molecular scale, early proposed auxetic polymers comprised covalently-bonded molecular honeycomb structures based on the macroscale re-entrant hexagonal honeycomb geometry known to realize auxetic response (Evans et al. 1991). Molecular modeling has shown that such molecular honeycombs would indeed be auxetic, but their heavily cross-linked structure renders them largely intractable from a synthesis point of view. Other theoretical molecular-level auxetic polymers have included 2D molecular honeycombs based on alternative known macroscale auxetic mechanisms (e.g. cooperatively rotating triangles) (Grima et al. 2005), 3D 'twisted chain' auxetics having a coupled polydiacetylene chain network (Baughman and Galvão 1993) and site-connectivity driven main-chain liquid crystalline polymers (He et al. 1998). In this latter example, the auxetic effect is achieved through chain separation driven by stress-induced rotation of laterally attached rods in main chains comprising rigid rod units connected by flexible spacer units. However, a fully developed synthetic molecular-level auxetic polymer remains elusive.

The reports of auxetic behavior in crystalline cellulose, therefore, provide natural systems to investigate in the drive to identify structures and mechanisms leading to auxetic response in crystalline polymers at the molecular scale. Conversely, the aforementioned reports in the literature of single-crystal mechanisms for auxetic response in polymers, and indeed other crystalline systems such as some silicates (Yeganeh-Haeri et al. 1992; Keskar and Chelikowsky 1992; Alderson and Evans 2002) and zeolites (Grima et al. 2000), suggest an investigation into a possible single-crystal explanation for the auxetic response in cellulose I_{β} is required.

In terms of crystal structure, cellulose I_{β} has a monoclinic unit cell within which there are two cellobiose chains aligned parallel to the crystallographic *c* axis (Nishiyama et al. 2002; French 2014). Matthews et al. (2006) note that Nishiyama's structure is in good agreement with the structure derived earlier by Finkenstadt and Millane (1998). The cellobiose repeat units are bonded by covalent bonds along each chain. One chain (the origin or corner chain) passes through atomic coordinates (0, 0) and the other chain (the center chain) passes through (1/2, 1/2) in the *a*- b plane (see Fig. 1a). The center chain is translated along c by $\sim c/4$ with respect to the corner chain. Each neighboring chain is bonded by hydrogen bonding to form stable planar sheets in the b-c plane, and neighboring sheets are held together by weak van der Waals-type interactions and weak C-H···O bonds to form a stacked sheet structure along the a axis (Simon et al. 1988). The precise nature of the packing of the chains in cellulose allomorphs has been the subject of much investigation over many years. Alternative packing arrangements of the chains are present in the literature and distinct chains can rotate about the c axis. A host of packing conformations are, then, possible and a number of these were identified in stable form in Molecular Dynamics simulations (Kroon-Batenburg et al. 1996). The chains pack in the 'parallel up' configuration in cellulose I_{β} (Nishiyama et al. 2002).

In kraft cooked Norway spruce, the kraft cooking process removes the lignin-hemicellulose matrix leading to a swelling of the crystallographic a lattice parameter (Peura et al. 2006). Swelling of the lattice parameter a is also known to occur during dehydration of crystalline cellulose (Zabler et al. 2010).

A number of studies have been undertaken on the mechanical properties of cellulose. A Mori–Tanaka approach and self-consistent scheme have been employed to model the elastic properties of a distribution of cellulose I_β crystallites in an amorphous matrix (nanofibrillated cellulose), predicting a ~56 % loss of stiffness compared to that of cellulose crystals along the main axis (Josefsson et al. 2013). Considering purely crystalline cellulose I_β, commercial

bleached ramie fibre (Nakamura et al. 2004), and early- and late-wood and juvenile and mature wood of Norway spruce (Peura et al. 2006) possess positive crystalline cellulose I_{β} Poisson's ratios. These are consistent with ab initio density functional theory (DFT) calculations of crystalline cellulose I_{β} where the Poisson's ratio is calculated to vary between +0.003 and +0.715, depending on direction (Dri et al. 2013). The dramatic modification of the v_{31} Poisson's ratio from positive to negative value as a result of the kraft cooking process suggests a possible mechanistic link with the swelling of the *a* lattice parameter upon removal of the lignin-hemicellulose matrix.

Numerous values for the Young's modulus along the chain axis of crystalline cellulose have been reported, showing wide variation typically in the range 134–220 GPa (Sakurada et al. 1962; Tashiro and Kobayashi 1991; Salmén 2004; Diddens et al. 2008), and theoretical upper limits as high as 246 and 319 GPa (Gillis 1969). Transverse Young's modulus values of 15 and 27.2 GPa have also been reported (Salmén 2004; Diddens et al. 2008). The variability in reported Young's moduli values has been attributed in the DFT calculations of crystalline cellulose I_{β} by Dri et al. (2013) to misalignment of crystals displaying highly anisotropic elastic response.

Molecular modeling can provide insight not achievable experimentally, but for crystalline cellulose structure and properties is challenging due in part to the presence of strongly polar bonds in close proximity, and the large range of configurations and conformations that can be formed (Foley et al. 2012). This is complicated by the variability of experimental

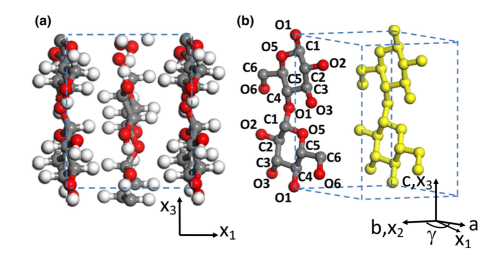


Fig. 1 Unit cell of crystalline cellulose I_{β}

data for validation, which may be related to microfibril history, size and aspect ratio (Matthews et al. 2012). Consequently, numerous carbohydrate-specific (Foley et al. 2012) and non-carbohydrate force fields have been used to model cellulose and related carbohydrate molecules. Carbohydrate force fields include the modified AMBER (Homans 1990), GLYCAM06 (Kirschner et al. 2008), CHARMM (Guvench et al. 2008) and GROMOS 45a4 (Lins and Hünenberger 2005) force fields. The CHARMM and AMBER-related GLYCAM06 force fields have, for example, recently been employed in Molecular Dynamics simulations and found to be reasonably accurate in predicting the crystal structures of a range of small molecule cellulose analogues (Miyamoto et al. 2016). CHARMM has also been used in Molecular Dynamics simulations of small microcrystallites of cellulose I_{β} in aqueous solution (Matthews et al. 2006). Unit-cell swelling (in particular the *a* lattice parameter), tilting of the molecular chains about the chain axis and inter-layer hydrogen bonds were predicted. Similar effects were predicted in hydrated cellulose I_{β} microfibrils with the GLYCAM06 force field, but not the Gromos 45a4 force field (Matthews et al. 2012). The unit-cell expansion and structures predicted by the CHARMM and GLYCAM06 force fields are similar to structures in simulations at elevated temperature (Matthews et al. 2011) and are consistent with experimental observation of anisotropic thermal expansion (Wada et al. 2010).

The CVFF (Consistent Valence Force Field) force field (Hagler et al. 1979a) is an example of one that is not recognized specifically for study of carbohydrates, but has been used to model the conformation of oligosaccharides (Hardy and Sarko 1993; Hagler et al. 1979b; Siebert et al. 1992). In a combined molecular mechanics and molecular dynamics study of the conformation of carbohydrate molecules, the CVFF force field was found to predict the conformation reasonably well in comparison to experimental nmr data (Asensio et al. 1995). A modified AMBER force field (Homans 1990), parameterized specifically for saccharides, was in qualitative agreement in the same investigation.

This paper reports a modeling investigation into the origins of the auxetic response in crystalline cellulose I_{β} . We aim to investigate qualitatively if there is a single-crystal mechanism that can explain the

experimental negative values of v_{31} following the kraft cooking process. Given the evident need for further experimental characterization to provide improved comparison with force field predictions, unresolved issues relating to the effects of microfibril size and shape on mechanical properties, and the sensitivity of predicted properties on carbohydrate force field parametrization, the intention here is not to achieve high quantitative accuracy in Poisson's ratio. Rather, we aim to probe the effects of unit-cell swelling, also observed following the kraft cooking process, on the structure and elastic properties of an infinite crystal of cellulose I_{β} . We undertake molecular mechanics energy minimizations using a range of force fields to identify a suitable one for the prediction of the structure of undeformed cellulose I_{β} . A more detailed analysis using the identified force field is then undertaken to identify the structure and elastic properties in the unloaded form and also with a stress condition applied to simulate the swelling along the a lattice parameter observed following the kraft cooking process.

Experimental section

Computational details

Molecular mechanics energy minimizations were carried out using the Cerius2 (version 4.6) and Materials Studio (version 4.1) molecular modeling packages (Accelrys). The following force fields, which have been parameterized to simulate the properties of organic networks, were assessed for their ability to accurately predict the structure of crystalline cellulose I_{β} : Compass (Sun 1998; Sun et al. 1998; Rigby et al. 1997), Dreiding (Mayo et al. 1990), Universal (Rappé et al. 1992), CVFF (Hagler et al. 1979a) and PCFF (polymer consistent force field) (Sun et al. 1994). These force fields have all been previously used to model cellulose, see for example Eichhorn and Davies (2006), Bazooyar et al. (2012), Dri et al. (2015), Asensio et al. (1995), and Mazeau and Heux (2003). The force field which gave the best overall prediction of lattice parameters and bond parameters (lengths and angles) was then selected for a more detailed investigation into the prediction of the elastic constants of single-crystal cellulose I_{β} before and after unit-cell swelling.

The model of cellulose I_{β} was built using the Visualiser module within Materials Studio. The monoclinic unit cell was constructed with cell parameters determined previously (Nishiyama et al. 2002) from X-ray and neutron diffraction: a = 7.784 Å, b = 8.201 Å, c = 10.380 Å and $\gamma = 96.55^{\circ}$. Space group P21 was assigned to the crystal. The carbon, hydrogen and oxygen atoms were then built into the cell according to the published fractional coordinates (Nishiyama et al. 2002). The refined crystal structure of cellulose I_{β} displays some hydrogen bond disorder (Nishiyama et al. 2002). In this work the hydrogens not included in the refined structure were assigned using the adjust hydrogen function in the materials visualizer module, which considers several factors such as common oxidation state, hydridization, formal charge, and number and order of bonds. A hydrogen bond cut-off of A-H = 3 Å was employed, close to the value of 2.8 Å used by Nishiyama et al. (2002) and Miyamoto et al. (2016). Periodic boundary conditions were applied to simulate an infinite cellulose I_{β} framework. Ewald summation was carried out to calculate the non-bond interactions (van der Waals and electrostatic). A convergence study showed the van der Waals energy varied significantly as the repulsive cut-off radius increased from 4 to 5 Å $(-33.77 \text{ to } -25.32 \text{ kcal mol}^{-1}, \text{ respectively})$, but did not change significantly as the cut-off radius was increased further $(-25.16 \text{ kcal mol}^{-1} \text{ at } 10 \text{ Å})$. Equally, the van der Waals energy varied between -16.10 and -25.21 kcal mol⁻¹ as the attractive cutoff radius increased from 6 to 12 Å and then remained virtually unchanged for further increases $(-25.24 \text{ kcal mol}^{-1} \text{ at } 18 \text{ Å})$. Accordingly, a cut-off radius of 15.5 Å was employed for the non-bond interactions, corresponding to the Fine quality setting in Materials Studio. For comparison, Miyamoto et al. (2016) employed switching functions from 12.0 to 14.0 Å for van der Waals interactions, and a 12 Å cut off for electrostatic interactions in simulations of the crystal structures of a range of small molecule cellulose analogues. Matthews et al. (2006) employed a non-bond interaction cut-off of 15 Å, and Matthews et al. (2012) applied a non-bonded cut-off distance of 10 Å for simulations using the CHARMM35 and GLYCAM06 force fields, whereas they used a longrange electrostatic interactions cutoff of 8 Å and a twin-range cut-off (inner cut-off of 8 Å and outer cutoff of 14 Å) for van der Waals interactions for the Gromos 45a4 force field.

The potential energy of the undeformed structure was minimized using the smart minimizer option, combining the steepest descent, ABNR (adjusted basis set Newton–Raphson) and quasi-Newton methods. The quality of the geometry optimization calculation was set to Fine within *Materials Studio* 4.1, corresponding to an energy convergence criterion of less than 0.0001 kcal mol⁻¹. During minimization, the P21 space group symmetry was converted to P1 and all cell parameters were set as variables. Convergence was achieved within the maximum number of minimization steps which was set at 5000 iterations.

The second derivative of the energy expression was used to compute the stiffness matrix *C*, and the on-axis elastic constants were obtained directly from the compliance matrix. The on-axis Poisson's ratios (v_{ij}) and Young's moduli (E_i) were calculated from the compliance coefficients (s_{ij}) using:

$$v_{ij} = -\frac{s_{ji}}{s_{ii}} \tag{1}$$

$$E_i = \frac{1}{s_{ii}} \tag{2}$$

where the subscripts *i* and *j* refer to directions x_i and x_j parallel and perpendicular to the loading direction, respectively. The mutually orthogonal principal axes x_2 and x_3 used for the description of mechanical properties in *Cerius*² and *Materials Studio* coincide with the *b* and *c* lattice parameter crystallographic directions, respectively, and the *a* lattice parameter crystallographic direction lies in the plane of the x_1 and x_2 principal axes but at an angle γ to the x_2 axis—see Fig. 1b.

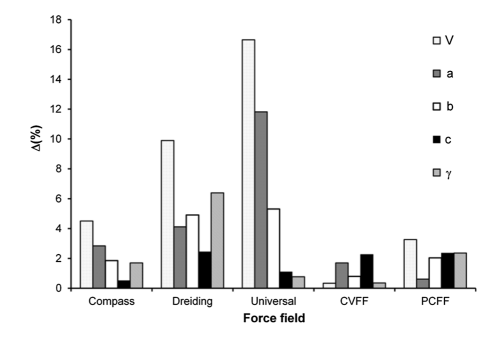
Results

Force field selection

To assess the variation between predicted and experimental values for each force field, a discrepancy factor was defined by:

$$\Delta = \frac{|x_{\exp} - x_{\text{mod}}|}{x_{\exp}} \times 100\%$$
(3)

Fig. 2 Discrepancy factors for the force field predictions (*Materials Studio*) of unit-cell parameters compared with the experimental data of Nishiyama et al. (2002)



where x = a, b, c, γ , bond angle, volume, etc. and the subscripts *exp* and *mod* represent experimental and predicted model values, respectively.

Unit-cell parameters

The predicted lattice parameters and unit-cell volume (*V*) of cellulose I_{β} for all force fields employed are compared with published experimental data (Nishiyama et al. 2002) in the Supporting Information (Online Resource Table 1) and the calculated discrepancy factors are shown in Fig. 2.

The CVFF force field gives the closest agreement with the experimental data, generally agreeing to within approximately 2 % or better for the lattice parameters and volume. The Universal force field fares the worst of the five force fields studied in this respect, typically being an order of magnitude worse in agreement than the CVFF force field. The CVFF force field has been parameterized using peptide and protein structures and has been extended to handle more general systems having similar functional groups, whereas the Universal force field is a general purpose force field with full coverage of the periodic table. The Universal force field is known to have only moderate accuracy for the prediction of geometries and conformational energy differences of organic molecules.

Bond parameters

All force fields reproduced the experimental covalent bond lengths to within better than 3 % discrepancy, with the exception of the C4–C5 bond length in the corner chain where the discrepancy was <4.5 % (Fig. 3 and Supporting Information Online Resource Table 2). The discrepancies between the force field model predictions and the experimental data (Nishiyama et al. 2002) for selected torsion angles are shown in Fig. 4. With the notable exception of the hydroxymethyl group torsion angle χ' (C4–C5–C6– O6) for the center chain, the force fields generally predict the torsion angles for the center and corner chains to better than 10 % discrepancy. The force fields predict χ and χ' values tending towards those for an ideal tg conformation (180° , -60°) whereas the experimental values for the center chain in particular deviate from ideality ($\chi = 158^{\circ}$ and $\chi' = -83^{\circ}$)— Supporting Information Online Resource Table 3. The CVFF force field appears to reproduce the torsion angles of the corner chain better than any of the other force fields. There is no clear cut best force field for the prediction of the torsion angles of the center chain.

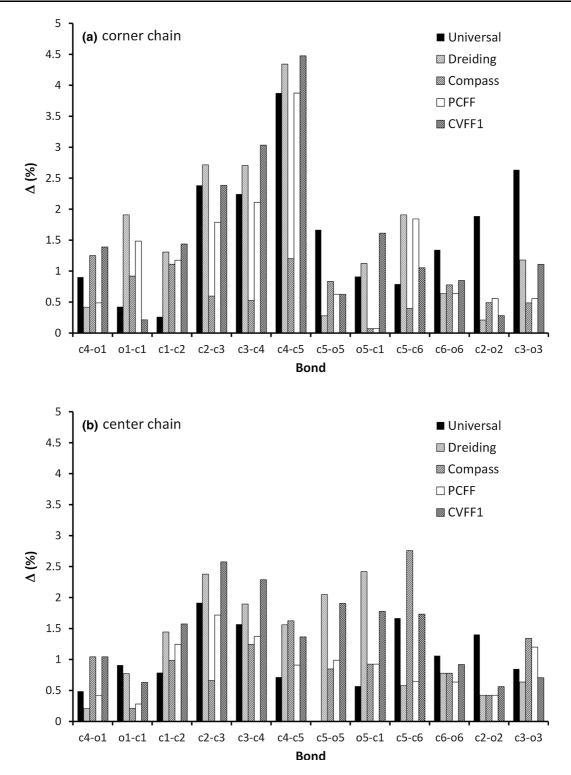
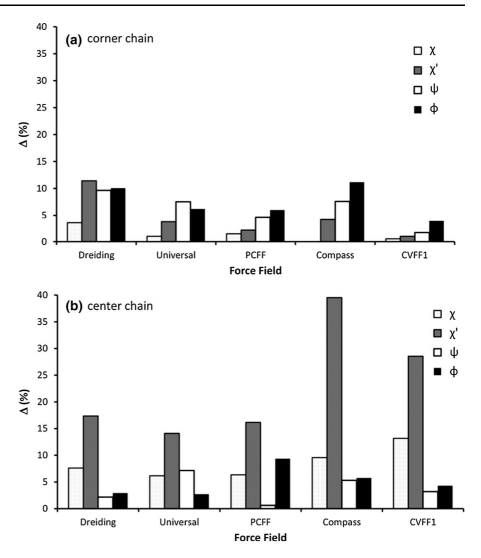


Fig. 3 Discrepancy factors for the force field bond length predictions compared with the experimental data of Nishiyama et al. (2002)

Fig. 4 Discrepancy factors for the force field bond angle predictions compared with the experimental data of Nishiyama et al. (2002)



Based on the above modeling of the structural parameters of cellulose I_{β} , the CVFF force field was selected for a more detailed investigation into the structure and mechanical properties of cellulose I_{β} .

Detailed undeformed structure investigation

Three models employing the CVFF force field have been employed in this part of the investigation. CVFF1 is the model developed above. CVFF2 incorporated a triaxial stress condition in order to reproduce the undeformed experimental lattice parameters for cellulose I_{β}. Finally, CVFF3 incorporated a stress in the x_1 direction to simulate the swelling along this axis observed during processing in the experimental measurement of negative Poisson's ratio response in kraft

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cooked Norway spruce (Peura et al. 2006). The triaxial stress applied in the CVFF2 model was determined from:

$$\begin{cases} \sigma_1 \\ \sigma_2 \\ \sigma_3 \end{cases} = \begin{cases} c_{11} & c_{12} & c_{13} \\ c_{21} & c_{22} & c_{23} \\ c_{31} & c_{32} & c_{33} \end{cases} \begin{cases} \varepsilon_a \\ \varepsilon_b \\ \varepsilon_c \end{cases}$$
(4)

where:

$$\varepsilon_c = \frac{c_{\exp} - c_{\rm CVFF1}}{c_{\rm CVFF1}} \tag{5}$$

and c_{ij} are elastic stiffness constants (*i*, *j* = 1, 2, 3) predicted from CVFF1. ε_c is the strain in the *c* lattice parameter required to convert the lattice parameter from CVFF1 to the experimental value. The subscripts *exp* and CVFF1 represent experimental and CVFF1 model

predicted values, respectively. Similar expressions were used for ε_a and ε_b . The stresses calculated in this way and employed in the CVFF2 model were: $\sigma_I = 0.38$ GPa, $\sigma_2 = 0.32$ GPa and $\sigma_3 = -3.98$ GPa.

For the CVFF3 model, a preliminary assessment of the effect of a (swelling) stress in the x_1 direction showed the appearance of negative on-axis Poisson's ratios in the x_1 - x_3 plane occurred when $\sigma_I > 0.76$ GPa (Fig. 5). Hence a tensile stress along the x_I direction of $\sigma_I = 0.8$ GPa was applied to the CVFF1 model to simulate both the swelling of lattice parameter *a* and the appearance of auxetic behavior observed after processing by Peura et al. (2006).

The unit-cell parameters and bond torsion angles for the CVFF1, CVFF2 and CVFF3 models are presented in Table 1. As expected, the CVFF2 model is in very close agreement with experimental unit-cell parameters (better than 0.3 % discrepancy for all parameters). The CVFF3 model displays the intended swelling in lattice parameter a, showing an increase of 7 % over the CVFF1 value. This is accompanied by a decrease in b by almost 3 % relative to CVFF1. Lattice parameter c remains largely unchanged between CVFF1 and CVFF3. There is little significant variation between the 3 models in the prediction of the torsion angles.

The hydrogen bond disorder reported experimentally for cellulose I_{β} (Nishiyama et al. 2002) makes an accurate prediction of the hydrogen bond network very difficult for force field-based energy minimizations of the type reported in this paper. Nevertheless,

Table 1 Comparison of the CVFF1, CVFF2 and CVFF3 model unit-cell parameter and bond torsion angle predictions with the experimental data of Nishiyama et al. (2002)

	CVFF1	CVFF2	CVFF3	Expt
a (Å)	7.652	7.787	8.188	7.784
b (Å)	8.136	8.201	7.898	8.201
c (Å)	10.615	10.388	10.607	10.380
γ (°)	96.8	96.2	95.1	96.5
V (Å ³)	656.1	659.5	683.2	658.3
Corner ch	nain			
χ (°)	169	170	172	170
χ′ (°)	-69	-70	-67	-70
ψ (°)	-144.7	-146.4	-143.6	-142.3
φ (°)	-94.7	-92.2	-94.5	-98.5
Center ch	ain			
χ (°)	179	179	180	158
χ' (°)	-59	-61	-59	-83
ψ (°)	-142.4	-143.6	-142.6	-147.1
φ (°)	-92.5	-90.2	-91.8	-88.7

we report the hydrogen bonding predicted from the CVFF1 model in Table 2 and Online Resource Fig. 1.

The hydrogen bonding network is predicted to comprise of O2–H···O6, O3–H···O5 and O6–H···O3 in the *b*–*c* plane for both the corner and center sheets. An additional O2–H···O1 hydrogen bond is predicted for the corner sheet. No hydrogen bonding is predicted between neighboring sheets by the CVFF1 model. Given the complexity of the multiple hydrogen bond network in cellulose I_{β} , the level of agreement

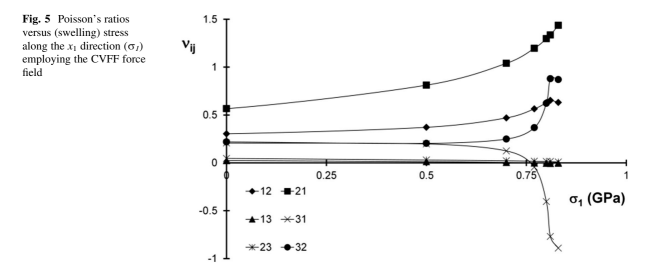


Table 2Comparison of theCVFF1 model hydrogenbonding parameters withthe experimental data ofNishiyama et al. 2002

D–H···A	d(D-H)	d(D–H) (Å)		d(H…A) (Å)		<dha (°)<="" th=""><th colspan="2">$d(D \cdots A)$ (Å)</th></dha>		$d(D \cdots A)$ (Å)	
	Exp	CVFF1	Exp	CVFF1	Exp	CVFF1	Exp	CVFF1	
Corner chain	1								
O2-H…O6	0.977	0.971	1.832	1.863	158.72	165	2.765	2.786	
O2−H…O1	0.977	0.971	2.285	2.378	110.28	109.2	2.799	2.848	
O3–H…O5	0.979	0.972	1.966	1.762	137.08	168.6	2.767	2.722	
06–H…O3	0.979	0.974	2.04	1.787	144.26	153.5	2.892	2.695	
Center chain									
O2-H···O6	0.982	0.973	1.904	1.73	165.12	158.4	2.865	2.659	
O3–H…O5	0.983	0.972	1.752	1.754	162.23	173.5	2.705	2.722	
O6–H…O3	0.985	0.971	1.779	1.921	156.61	158.3	2.711	2.85	

between the predicted CVFF1 model hydrogen bond parameters and the values proposed from X-ray and neutron scattering studies (Nishiyama et al. 2002; Table 2) is considered acceptable. With the exception of additional C4–H···O2 bonds between adjacent sheets predicted by the CVFF3 model there are no significant additional or different features predicted from the CVFF2 and CVFF3 models.

Projections of the unit cell in the x_1 - x_2 plane, i.e. normal to the chain axes, are shown in Fig. 6 for the experimental and minimized CVFF1 and CVFF3 structures. The experimental atomic coordinates produce corner and center chains having similar rotations (tilt angles δ_1 and δ_2 , respectively) about the *c* axis. In the CVFF1 model the corner and center chains undergo anticlockwise and clockwise rotation, respectively, relative to the experimental chain orientation. The opposing rotations about the *c* axis of the two distinct chains in the model structures are exaggerated further in the CVFF3 model, and are accompanied by the presence of the aforementioned C4–H…O2 bonds between adjacent sheets and the intended swelling of the unit-cell along the x₁ direction. Detailed elastic constants investigation

The compliance coefficients predicted from the CVFF1, CVFF2 and CVFF3 models, calculated from the 2nd derivative of the energy expression, are presented in the Online Resource Table 4. Table 3 contains the values of v_{ij} , and E_i calculated from the compliance coefficients using Eqs. (1) and (2).

The Young's modulus along the chain length (E_3) is higher for CVFF2 than for CVFF1, whilst the two transverse Young's moduli are lower. However, the CVFF1 and CVFF2 models return very similar positive Poisson's ratios to each other, generally agreeing to two decimal places for each Poisson's ratio, implying the triaxial stress applied in CVFF2 to correct for deviation of the CVFF1 lattice parameters from the known experimental values has little effect on the predicted Poisson's ratio response.

The CVFF3 model predicts a slightly lower E_3 and order of magnitude lower E_1 and E_2 values than the CVFF1 and CVFF2 models. Higher positive on-axis Poisson's ratios in the x_1 - x_2 plane are predicted by the CVFF3 model. Positive on-axis Poisson's ratios are

Fig. 6 Projected structures of cellulose I_{β} in the x_1 - x_2 plane: **a** experimental, **b** minimized CVFF1 model (including O6–H···O3 interchain hydrogen bonds), and **c** minimized CVFF3 model (including C4–H···O2 intersheet bonds)

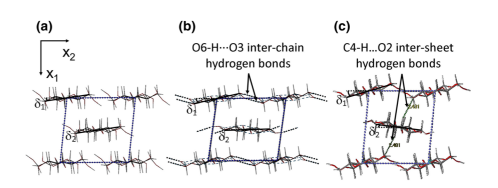


Table 3 Poisson's ratios and Young's moduli (GPa) for cellulose I_{β} predicted using the 2nd derivative of the energy expression in the CVFF1, CVFF2 and CVFF3 models

	CVFF1	CVFF2	CVFF3
v ₁₂	0.303	0.305	0.629
v_{13}	0.023	0.018	-0.005
v_{21}	0.566	0.569	1.297
V ₂₃	0.046	0.039	0.015
v_{31}	0.207	0.207	-0.406
V ₃₂	0.219	0.240	0.623
E_1	21.0	17.4	2.1
E_2	39.1	32.4	4.3
E_3	186.0	201.5	179.6

also predicted in the x_2-x_3 plane, with v_{23} and v_{32} being, respectively, lower and higher than predicted by the CVFF1 and CVFF2 models. As noted above, negative on-axis Poisson's ratios are predicted by the CVFF3 model in the x_1-x_3 plane, reaching as low as $v_{31} = -0.406$.

To check the auxetic response in the x_1 - x_3 plane for the CVFF3 model, energy minimizations were performed for 0.5 GPa increments of stress along the x_3 axis, maintaining $\sigma_1 = 0.8$ GPa and $\sigma_2 = 0$ GPa in all cases. The choice of stress increment along x_3 is similar to that used in previous molecular mechanics studies on the response of crystalline polymers to an applied stress (Evans et al. 1995; Nkansah et al. 1994; Alderson et al. 2005). Figure 7 shows the calculated true strains in the transverse x_1 and x_2 directions as a function of strain in the loading x_3 direction. The true strains were calculated from the lattice parameters as follows:

$$\varepsilon_1 = \ln\left(\frac{a\sin\gamma}{a_0\sin\gamma_0}\right), \ \varepsilon_2 = \ln\left(\frac{b}{b_0}\right), \ \varepsilon_3 = \ln\left(\frac{c}{c_0}\right)$$
(6)

where the subscript '0' indicates the parameter value at $\sigma_3 = 0$ GPa. The strains along x_1 and x_2 increase and decrease, respectively, with increasing strain along x_3 , consistent with the negative and positive signs of v_{31} and v_{32} calculated from the 2nd derivative of the energy expression (Table 3). The Poisson's ratios in the undeformed CVFF3 model are determined from the negative of the slope of the 2nd order polynomial fit curve in Fig. 7 at $\varepsilon_3 = 0$ for each case, giving $v_{31} = -0.559$ and $v_{32} = +0.718$. These compare to values calculated from the 2nd derivative of the energy expression (Table 3) of $v_{31} = -0.406$ and $v_{32} = +0.623$.

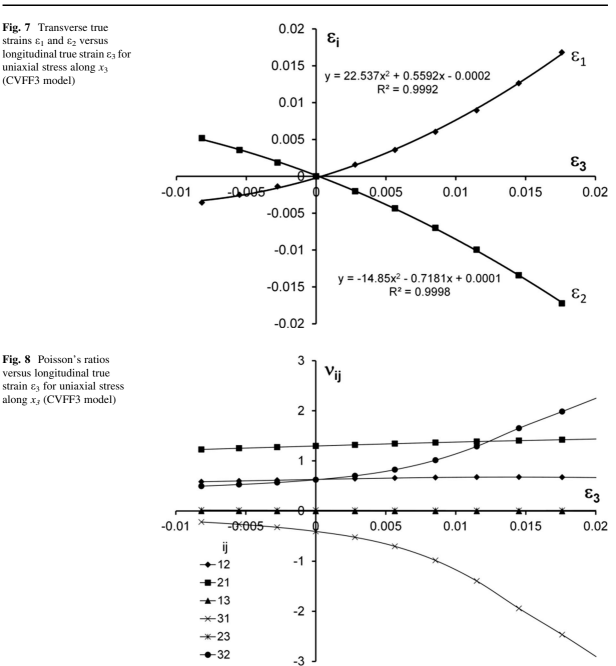
Figure 8 shows the CVFF3 model Poisson's ratios (2nd derivative method) as a function of applied true strain for loading along the chain (x_3) direction. The negative and positive Poisson's ratios v_{31} and v_{32} , respectively, are both predicted to vary with strain, with magnitude increasing with increasing strain. Poisson's ratios v_{12} , v_{21} , v_{13} and v_{23} are predicted to be almost independent of strain by comparison. Preliminary energy minimizations (data not shown) for the CVFF1 and CVFF2 models showed these two models to display identical trends to each other with all Poisson's ratios remaining constant (and positive) over the strain range covered in Fig. 8.

Structural evolution with stress along the x_3 direction

Given that the CVFF3 model predicts the on-axis auxetic behavior of cellulose I_{β} , this model was selected to perform a detailed investigation of the variation in the structure of cellulose I_{β} under loading in the x_3 direction, in an attempt to identify the mechanism for auxetic response.

Chain unfolding in the x_2 – x_3 *plane*

Figure 9a shows a repeat cellobiose unit with best-fit plane ABDE created by Materials Studio shown as a semi-transparent surface with an arrow normal to the plane indicating its orientation. The best fit plane is defined as a plane where the (mass weighted) root mean square distances from the plane to the selected atoms are minimized. The selected atoms used to generate the best-fit plane ABDE were O1, C1, C2, C3, C4, C5, C6, O2, O3, O5, O6 and O1(O4). For clarity, only atoms related to the torsion angles ϕ and ψ , and the bridging angle τ (=C1–O1–C4) are labelled in Fig. 9a. Plane ABDE is almost parallel to the x_2 - x_3 plane, indicated by the arrow normal to the plane being aligned nearly parallel to the x_1 axis. Atom O1 lies above the plane ABDE and O4 below the plane ABDE (i.e. bonds C1–O1 and O4–C4 are over and below the plane ABDE, respectively). The glycosidic torsion angles ψ and ϕ are defined as the angle between plane C5-C4-O4(O1) and plane C4-O4(O1)-C1 and the



angle between planes O5–C1–O1 and C1–O1–C4, respectively.

The glycosidic bridging angle τ is predicted to increase with increasing tensile strain in the loading (x_3) direction (Fig. 9b), indicating an unfolding of the backbone chain along the x_3 axis in the x_2 - x_3 plane. The chain unfolding in the x_2 - x_3 plane is accompanied by a predicted increase with loading strain in the O1…O4(O1) distance (Fig. 9c) which implies some

stretching of the cellobiose unit along the axis connecting O1 and O4(O1) also occurs.

Chain rotation in the x_1 – x_2 *plane*

Figure 9b also shows the predicted variations of the helical parameters ϕ (C4–O1–C1–O5) and ψ (C1–O4(O1)–C4–C5). ϕ is predicted to increase and ψ is predicted to decrease with increasing tensile strain in

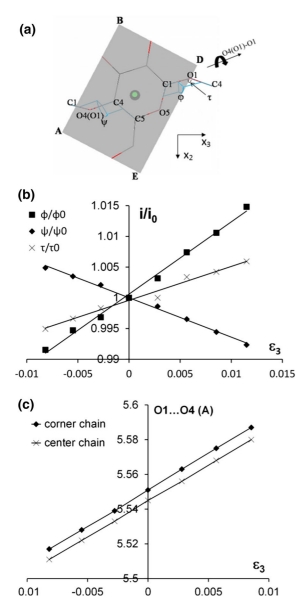


Fig. 9 a Projection in the x_2-x_3 plane of a repeat cellobiose unit with best-fit plane ABDE created by *Materials Studio*. ϕ and ψ are the torsion angles defined by (C4–O1–C1–O5) and (C1– O4(O1)–C4–C5), respectively. τ is angle C1–O1–C4. **b** Predicted change in bridging angle τ and torsion angles ϕ and ψ ($i = \tau$, ϕ or ψ) as a function of strain in the loading (x_3) direction. **c** Predicted change in O1…O4(O1) distance in cellobiose unit as a function of strain in the loading (x_3) direction

the loading (x_3) direction. This corresponds to the plane ABDE rotating about the O1–O4(O1) axis. Since the O1–O4(O1) axes in adjacent cellobiose units are aligned at angles of equal but opposite sign with respect to the x_3 axis, the rotation of plane ABDE

about the O1–O4(O1) axis corresponds to a rotation of the helical chain about the helical (x_3) axis.

This is shown in Fig. 10a which shows that the cellulose chains appear as tilted 'rod' segments (terminated by C6 atoms) interconnected by hydrogen bonds when projected onto the x_1 - x_2 plane. Tilt angles δ_1 and δ_2 , previously referred to in Fig. 6, are defined by the angle the cellulose 'rods' in the corner and center chains make with the x_2 axis in the x_1 - x_2 projection, respectively (positive values correspond to anti-clockwise rotation). The CVFF3 model predicts values for the undeformed structure of $\delta_1 = 17.2^\circ$ and $\delta_2 = -6.7^\circ$. The magnitudes of δ_1 and δ_2 are predicted to increase with increasing tensile strain along the x_3 direction (Fig. 10b). In other words, each cellulose chain is predicted to rotate about its axis in response to a uniaxial load applied along x_3 .

Cellulose chains in adjacent sheets are predicted to be connected by C4–H···O2 bonds (Fig. 6c), whereas inter-chain bonding within a sheet occurs via O6-H…O3 hydrogen bonds (Fig. 6b). This is consistent with previous reports showing no hint of inter-sheet O-H…O hydrogen bonds, and cellulose sheets held together by C-H···O bonds and hydrophobic interactions (Finkenstadt and Millane 1998; Nishiyama et al. 2002; Claffey and Blackwell 1976). The length between the (donor) C4-H in the center chain and O2 (accepter) in the corner chain is predicted to remain constant at 2.491 Å as stress is applied along the x_3 direction (Table 4). A slight decrease in the distance between the donor hydrogen and acceptor oxygen is predicted with increasing applied stress for the intra-sheet O6–H···O3 hydrogen bond (d(H···A) decreases from 2.643 to 2.631 Å as stress increases in the range $-2 < \sigma_3 < +2$ GPa). The C6…C6 distances defining the rod segments in Fig. 9a increase slightly with increasing tensile stress along x_3 for both the corner and center chains, consistent with the predicted increase in glycosidic bridging angle τ .

Chain unfolding in the x_1 *–* x_3 *plane*

The projection of the cellulose I_{β} structure in the x_1 - x_3 plane is presented in Fig. 11a, showing the covalently bonded chains aligned along the x_3 axis. The bridging O1 atoms are labeled for clarity. The projected structure can be characterized by a zigzag framework connecting the O1 bridging atoms along each chain. The angle the projected O1...O4(O1) distance makes

σ_3 (GPa)	C4–HO2 d(D…A) (Å)	O6–HO3 d(D…A) (Å)	Corner chain C6…C6 (Å)	Center chain C6…C6 (Å)
2	2.491	2.631	7.674	7.643
1.5	2.491	2.632	7.665	7.62
1	2.491	2.633	7.656	7.608
0.5	2.491	2.634	7.646	7.597
0	2.491	2.636	7.637	7.585
-0.5	2.491	2.637	7.627	7.574
-1	2.491	2.639	7.618	7.562
-1.5	2.491	2.641	7.608	7.551
-2	2.491	2.643	7.597	7.54

Table 4Predictedhydrogen bond andcellobiose chain distancesagainst applied stress in the x_3 direction (CVFF3 model)

with the x_3 axis in the x_1-x_3 plane (defined as α_1 and α_2 for the corner and center chains, respectively— Fig. 11a) is predicted to decrease with increasing tensile strain in the loading (x_3) direction for both the corner and center chains, as shown in Fig. 11b. The cellulose chains are, then, also predicted to unfold along x_3 in the x_1-x_3 plane in response to a uniaxial tensile load applied along x_3 .

Discussion

In this work, molecular mechanics has been used to predict the structure and elastic properties of singlecrystal cellulose I_{β} . The CVFF force field has been found to reproduce the lattice parameters, bond structure and hydrogen bond structure reasonably well. In the undeformed state the CVFF force field predicts positive on-axis Poisson's ratios. The on-axis Poisson's ratios are all predicted to be positive and remain virtually unchanged when a triaxial stress is applied in order for the predicted lattice parameters to reproduce exactly the experimentally determined values. The positive Poisson's ratios are consistent with reported values for crystalline cellulose from normally grown wood, which have been attributed to the presence of the amorphous lignin-hemicellulose matrix (Peura et al. 2006).

Applying a stress along the x_1 direction, to simulate swelling along this direction, leads to the predicted appearance of negative values of the major Poisson's ratios in the x_1 - x_3 plane. This is consistent with the previous experimental measurement of a negative v_{31} Poisson's ratio in kraft cooked Norway spruce, in which the removal of lignin and hemicelluloses during cooking leads to the swelling of the crystallites in the x_1 direction (Peura et al. 2006). The values of $v_{31} = -0.559$ and -0.406 from the slope of the transverse strain vs axial strain curve and the 2nd derivative of the energy expression (Table 3), respectively, compare with the experimental average values of v_{31} in the ranges -0.26 to -1.17 (before yield point) and -0.86 to -1.05 (after yield point) (Peura et al. 2006). The results suggest an alternative or additional mechanism for auxetic behavior to the microfibril non-alignment and cell-cell wall layer shearing interactions proposed by Peura et al. (2006).

The swelling simulation in CVFF3 employed a stress along the x_1 direction to simulate swelling since it is expected that the inter-plane direction is most affected by removal of the hemicellulose-lignin matrix and presence/absence of moisture (Zabler et al. 2010). On the face of it the employed stress of 0.8 GPa is a very large stress to use since the theoretical tensile strength of crystalline cellulose is ~2 GPa along the chain (x_3) direction (Mark 1967), for example. However, 0.8 GPa is within the range of stress magnitudes employed in the CVFF2 model to bring the CVFF force field unit-cell length predictions in line with experiment. Hence caution should be applied when considering the absolute value of stress employed in the swelling simulation.

The amount of swelling due to the kraft cooking process is not clear from Peura et al. (2006). For comparison, a 0.6 % expansion in *a* during dehydration of crystalline cellulose has been reported (Zabler et al. 2010), and a 5 % expansion in *a* occurs on heating cellulose I_{β} to the transition to high temperature cellulose I (Wada et al. 2010). The expansion in the CVFF3 simulation (relative to the CVFF1 simulation) reported here (Table 1) is 7 %. In this work we have been concerned with the effects of swelling on

the mechanical properties of cellulose I_{β} . More sophisticated modelling treatments for simulating swelling than those presented here have been reported including, for example, molecular dynamics simulations of small microcrystallites of cellulose I_{β} in aqueous solution (Matthews et al. 2006, 2012).

The CVFF1 and CVFF2 Young's moduli (Table 3) are in agreement with previous axial and transverse Young's moduli ranges from the literature of 134–220 GPa (Sakurada et al. 1962; Gillis 1969; Tashiro and Kobayashi 1991; Salmén 2004; Diddens et al. 2008) and 15–27.2 GPa (Salmén 2004; Diddens et al. 2008), respectively. The CVFF3 energy minimizations indicate the effect of swelling of the *a* lattice parameter has little effect on the Young's modulus in the chain direction (E_3) but reduces the transverse Young's moduli (by an order of magnitude for the amount of swelling considered in the models).

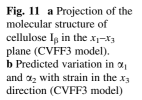
The mechanism responsible for the negative v_{31} response has been explored by examining the evolution of the crystallographic structure in energy minimizations employing a uniaxial stress in the x_3 direction. The cellobiose chains are predicted to undergo rotation in the x_1-x_2 plane (transverse to chain axis and loading (x_3) direction—Fig. 10). The C4-H...O2 inter-sheet interaction distance was found to remain constant with applied stress along the x_3 direction and, therefore, effectively acts as a rigid spacer unit between the rotating cellobiose chains. This enables the development of a conceptual model providing a mechanistic explanation for the auxetic response. In the conceptual model the projection of the structure in the x_1-x_2 plane is simplified in a first approximation to an array of rotatable rigid rods (projected cellobiose chains) connected by a series of intra-layer and inter-layer inextensible spacers (projected O-H···O and C4-H···O2 interaction distances, respectively). This is shown in Fig. 12 where the rigid spacers corresponding to the projected O-H...O intralayer distances connect adjacent rigid rods in a cornerto-corner manner [one bottom corner to one top corner in the untilted ($\delta = 0$) configuration (Fig. 12b)]. The projected C4-H···O2 rigid spacers are connected to the 'corner sheet' rods at a corner adjacent to the connected 'center sheet', and to the 'center sheet' rods at a fixed location on the edge facing the connected 'corner sheet'. The untilted configuration shown in Fig. 12b possesses a rectangular unit-cell in the $x_1 - x_2$ plane.

If it is assumed that the angle the inter-sheet rigid spacer makes with the center sheet rigid rod (assumed perpendicular in Fig. 12) remains constant irrespective of the tilt (rotation) of the rigid rods, then cooperative rotation of the rigid rod units by angle δ with respect to the x_2 axis leads to the structure shown in Fig. 12c, with a concomitant shearing of the unit cell in the x_1 - x_2 plane. This is consistent with the monoclinic unit-cell and tilted chain orientations predicted by the undeformed CVFF3 model (Fig. 12a). Increasing the rigid rod rotation (tilt angle δ) further causes an increase in lattice parameter *a* and a decrease in lattice parameter b and, therefore, expansion and contraction of the structure along the x_1 and x_2 directions, respectively (Fig. 12d). Figure 10 shows that such rigid rod rotation occurs in response to a tensile stress applied in the x_3 direction perpendicular to the plane of the structure ($\sigma_3 > 0$), and so this corresponds to negative and positive values for v_{31} and v_{32} , respectively, as predicted in the force field-based energy minimizations (Table 3; Fig. 8). Similarly, applying a compressive load along the chain axis to the structure in Fig. 12c would tend to remove the rigid rod tilt (i.e. tend towards the structure in Fig. 12b) resulting in contraction and expansion of the structure along the x_1 and x_2 directions, respectively, consistent with the CVFF3 model predictions in Fig. 7.

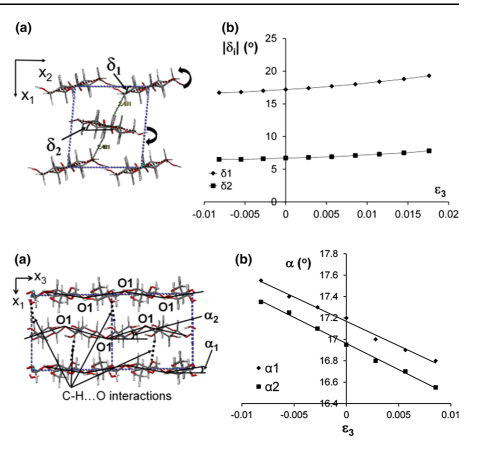
Clearly the 2D interpretation presented in Fig. 12 is a simplification of the cellulose I_{β} system. Whilst the force field energy minimizations indicate inter-sheet C4-H...O2 interactions act as rigid spacers in 3D (Table 4), a slight reorientation of this interaction length is also predicted such that the projected C4-H···O2 distance in the x_1 - x_2 plane increases slightly with increasing tensile stress along x_3 (the models indicate an increase in the projected length of $\sim 0.2 \%$ for an applied strain along x_3 of $\sim 1\%$). The projected C4–H···O2 distance is aligned most closely with the x_1 axis and so the increase in the projected length provides an increase in x_1 additional to that from $x_1 - x_2$ chain rotation, reinforcing the negative value of v_{31} . Similarly, the projected O-H…O intra-layer interaction distance, most closely aligned with the x_2 axis, decreases in the x_1 - x_2 plane with increasing tensile stress (by ~0.1 % for an applied strain of ~1 %), giving an additional decrease in x_2 to that from x_1 to x_2 chain rotation to reinforce the positive value of v_{32} .

The prediction of chain tilting and inter-layer hydrogen bonds due to swelling along the x_1 direction

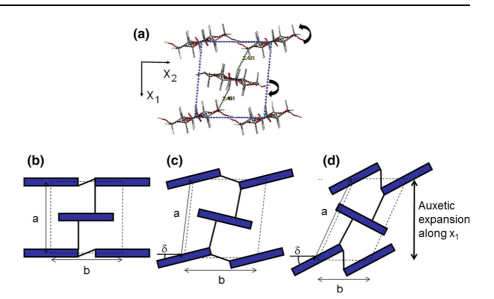
Fig. 10 a Projection of the structure of cellulose I_{β} in the x_1-x_2 plane, defining tilt angles δ_1 and δ_2 as the angle the C6...C6 length projected in the plane makes with the x_2 axis for the corner chain and center chain, respectively. b Predicted variation in the magnitudes of δ_1 and δ_2 with strain in the x_3 direction (CVFF3 model)



in an infinite crystal of cellulose I_{β} in this work has also been predicted in Molecular Dynamics simulations of small microcrystallites of cellulose I_{β} in aqueous solution using the CHARMM and GLYCAM force fields (Matthews et al. 2006; Matthews et al. 2012). Structures with tilts and rotating units have previously been identified as leading to auxetic behavior. Examples include rotation of tetrahedral sub-units in inorganic tetrahedral framework systems such as the α -cristobalite polymorph of crystalline silica (Yeganeh-Haeri et al. 1992; Alderson and Evans 2002), and in certain zeolitic structures (Grima et al. 2000). Rotating units have also been proposed in 'nodule-fibril' models for auxetic microporous polymers (Evans and Caddock 1989) and, at the molecular level, in auxetic liquid crystal polymers (He et al. 1998). In the polymer cases, the rotation of the units has been reported in the plane containing the loading direction. This work identifies rotation of tilted units in the plane transverse to the molecular chain axis and loading direction as a mechanism for auxetic response in crystalline polymeric systems.



The orientation of the cellobiose chain in the x_1-x_2 plane and the identification of the role of rotation of these chains about the x_3 -axis is analogous to the mechanism proposed for anisotropic (including negative) thermal expansion in pentacene crystals (Haas et al. 2007). The tilt angle identified in this work is analogous to the herringbone angle describing the packing of the assumed rigid pentacene molecules. With increasing temperature, the increase of the herringbone angle and the increasingly dominant vibrations about the long molecular axis cause a distinctly anisotropic expansion in the a-b plane, similar to that described in Fig. 12. The vibrating molecules at the corners of the pentacene unit cell interact with the one in the center, pushing each other further apart in one unit-cell direction as the vibration amplitude grows with temperature, with a concomitant contraction in the other lattice parameter giving rise to a negative coefficient of thermal expansion in this direction. In other words, the two principal directions in the x_1-x_2 plane undergo opposite expansion/contraction in response to thermal loading in pentacene **Fig. 12** a Projection of the CVFF3 model structure in the x_1 - x_2 plane. Conceptual model of a network of rotatable rigid rods interconnected by inextensible spacers mimicking the projected CVFF3 structure in the x_1 - x_2 plane: **b** no rotation of rigid rods, **c** intermediate rotation of the rigid rod units and **d** high rotation of the rigid rod units



crystals, whereas in cellulose I_β crystals this mechanism appears to be triggered by uniaxial mechanical loading along the x_3 direction.

Interestingly, cellulose I_{β} also displays anisotropic thermal expansion which becomes more dramatic upon phase transition to high temperature cellulose I (Wada et al. 2010). As noted above, high temperature cellulose I displays significant swelling of the *a* lattice parameter (relative to cellulose I_{β}) which undergoes expansion upon further heating, accompanied by a contraction of the b lattice parameter. These are the same responses as for a tension applied along the chain axis in the CVFF3 model presented here. Molecular dynamics simulations (CHARMM and GLYCAM06 force fields) predict the presence of tilted chains and inter-layer hydrogen bonds in high temperature cellulose I (Matthews et al. 2011). Taken together these suggest a mechanistic link between the thermal expansion and Poisson's ratio responses in high temperature cellulose I and kraft cooked cellulose I_{β} , respectively.

It is tempting to compare the projected zig-zag framework connecting the O1 bridging atoms along each chain and interconnecting C4–H···O2 inter-sheet interactions in the x_1 – x_3 plane (Fig. 11a) with a 2D reentrant hexagonal honeycomb structure. Deformation of a re-entrant hexagonal honeycomb by rotation of the ribs forming the honeycomb, corresponding to a reduction in honeycomb angle α (i.e. unfolding of the zig-zag structure along the x_3 direction) for a tensile stress in the x_3 direction, is known to produce a

negative value for v_{31} (Evans et al. 1995; Gibson et al. 1982; Masters and Evans 1996). The molecular mechanics energy minimizations presented here predict an unfolding of the molecular chains along the loading x_3 axis (Fig. 11b) consistent with this mechanism. However, the models also indicate the projection of the O1…O1 distance in the x_1-x_3 plane increases with increasing tensile stress along x_3 (projected distance increases by ~ 0.7 % for loading strain of ~ 0.9 %). This corresponds to stretching of the oblique ribs in a re-entrant hexagonal honeycomb, which is known to lead to a positive value of v_{31} (Masters and Evans 1996). From the analytical model expressions for a re-entrant hexagonal honeycomb (Masters and Evans 1996), the magnitude of change in projected O1…O1 length would appear to be sufficient to overcome the zig-zag unfolding mechanism driving the auxetic response.

This work has identified mechanisms of chain rotation and chain unfolding, and the crucial role of secondary bonding, in realizing auxetic response at the molecular chain level in a naturally-occurring macromolecular system. It is expected this will provide impetus to the further design and development of molecular-engineered synthetic auxetic polymers.

Conclusion

In this work we have attempted to interpret the structural evolution of the 3D structure of crystalline

cellulose I_{β} through consideration of key structural features in 2D projections. It is clear that the effects of bond stretching and reorientation of the 3D structure complicate an interpretation of the mechanisms responsible for auxetic behavior in such systems via a consideration of 2D projections. Nevertheless we have identified here two potential mechanisms operating in crystalline cellulose I_{β} : namely rotation of cellobiose chains transverse to the loading x_3 direction, and unfolding of the chains along the loading direction. The former mechanism appears to be the most likely of the two mechanisms to be responsible for the auxetic effect predicted by the molecular mechanics energy minimizations. In both mechanisms the presence of secondary (hydrogen) bonding between chains, perpendicular to the chain/loading direction, appears to be crucial in realizing auxetic behavior.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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