Microsecond kinetics in model single- and
double-stranded amylose polymers†

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Amylose, a component of starch with increasing biotechnological significance, is a linear glucose
polysaccharide that self-organizes into single- and double-helical assemblies. Starch granule packing,
gelation and inclusion-complex formation result from finely balanced macromolecular kinetics
that have eluded precise experimental quantification. Here, graphics processing unit (GPU) accelerated
multi-microsecond aqueous simulations are employed to explore conformational kinetics in model
single- and double-stranded amylose. The all-atom dynamics concur with prior X-ray and NMR data
while surprising and previously overlooked microsecond helix-coil, glycosidic linkage and pyranose
exchange are hypothesized. In a dodecasaccharide, single-helical collapse was correlated with linkages
and rings transitioning from their expected syn and C1 chair conformers. The associated microsecond
exchange rates were dependent on proximity to the termini and chain length (comparing hexa- and
trisaccharides), while kinetic features of dodecasaccharide linkage and ring flexing are proposed to be a
good model for polymers. Similar length double-helices were stable on microsecond timescales but the
parallel configuration was sturdier than the antiparallel equivalent. In both, tertiary organization restricted
local chain dynamics, implying that simulations of single amylose strands cannot be extrapolated to
dimers. Unbiased multi-microsecond simulations of amylose are proposed as a valuable route to
probing macromolecular kinetics in water, assessing the impact of chemical modifications on helical
stability and accelerating the development of new biotechnologies.

Introduction

Starch is a major plant energy storage material comprising a
source-dependent mix of two \( \alpha(1 \rightarrow 4) \) linked \( \alpha \)-glucose (Glc)
polysaccharides: linear amylose and \( \alpha(1 \rightarrow 6) \) branched amylo-
pectin. Amylose is smaller in size, typically less abundant and
pivotal to the aqueous insolubility and partial crystallinity of starch
granules. A major dietary component, amylose is also used as a
thickener, stabilizer and gelling agent in foods, cosmetics, textiles
and paper. It has recently been employed as a prototype biotechnol-
ological device, with notable successes in host-guest inclusion and
for improving the biocompatibility of carbon nanotubes, drugs and
flavor ingredients. These diverse current and potential applications
stem from the dynamic molecular shape of amylose, which gives
rise to characteristic helices and self-association.

A helical amylose structure was first reported in 1943. Subsequent X-ray studies have revealed three principal crystal
allomorphs: single-helical V-type amylose and the A- and B-type
double-helices (from cereals and tubers, respectively). Flexible
in solution, V-amylose has been found by X-ray analyses to include six, seven or eight monosaccharides per helical turn. The A and B forms are more rigid and differ only in their
packing arrangement, with six pyranoses per turn. Water plays a
key role in amylose shape and single-helix stability can be tuned by
solvent and chemical modifications that modulate inter- and
intra-molecular hydrogen-bonding (e.g., DMSO and alklylation). Interrupting these through-space interactions while maintaining
higher-order helical structures is a promising route to improving
the solubility of amylose-based materials. Precise quantification of aqueous conformational exchange
rates in amylose has eluded experimental techniques. For example,
iodine induced self-assembly rates can only be estimated to occur
in the sub-ms domain. Solution NMR has revealed single-
strand amylose to transition between compact coils and
extended helices but more detailed analyses using this techni-
que are restricted by severe resonance overlap.

The kinetics of conformational exchange in amylose glycosidic
linkages and pyranose rings also remain enigmatic. Reversible
rotation of the amylose \( \alpha(1 \rightarrow 4) \) glycosidic torsion through
\( \approx 180^\circ \), a syn → anti transition or band-flip, interrupts helical
shapes causing chain kinks that are associated with coiled struc-
tures in cyclic amyloses. Constituent rings may also be flexible

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in water and G1c conformational transitions are thought to underlie amylose elasticity.\textsuperscript{16} While the G1c \textsuperscript{4}C\textsubscript{1} chair is favored, the aqueous pyranose ring puckering equilibrium is sensitive to the chemical environment (e.g., substituents\textsuperscript{17} and anomeric configuration\textsuperscript{18}). Recent graphics processing unit (GPU) accelerated simulations have predicted \( \mu \)s conformational exchange to occur in \( \beta \)-D-glucose\textsuperscript{19} but this has not yet been explored in linear \( \alpha(1 \rightarrow 4) \) linked Glc chains.

Further conformational insights into aqueous amylose motions have been derived from computational studies of fragments using molecular dynamics\textsuperscript{20–23} and Monte Carlo\textsuperscript{24} methods. However, unbiased simulations have been limited by computational cost to short ns timescales and theoretical techniques that artificially enhance conformational sampling preclude calculation of accurate kinetic data for exchange between molecular shapes.\textsuperscript{25} While coarse grained modeling has provided macroscopic conformational predictions for large amylose chains,\textsuperscript{26} until recently\textsuperscript{27} such mesoscale approaches have neglected pyranose ring puckering and have also been based on predictions from ns simulations.

Here, GPU-accelerated all-atom simulations were used to explore the microsecond kinetics of amylose macromolecular motions in water, to improve understanding of biological function and to enable future engineering of derivative biotechnological materials. In particular, 10 \( \mu \)s simulations were carried out on a model amylose dodecasaccharide (1) and model dodecasaccharide double-helices, initiated in both antiparallel and parallel configurations (2 and 3). For comparison, 10 \( \mu \)s simulations of a constituent hexa-, tri- and monosaccharide (4, 5 and 6) were also performed (Fig. 1). These trajectories were used to identify and characterize \( \mu \)s conformational exchange rates for helix-coil, glycosidic linkage and pyranose ring transitions. Additionally, the effects of chain length, position, tertiary association and double-helix directionality on these degrees of freedom were investigated.

Results and discussion
Experimental consistency and microsecond exchange
The mean radius of gyration, \( R_g \), and the standard deviation of this mean (see Methods) in the simulations of 1, 2 and 3 (see Fig. 1 for structures) were 11.5 (\( \pm 1.1 \)) Å, 13.2 (\( \pm 0.3 \)) Å and 14.4 (\( \pm 0.1 \)) Å, respectively, consistent with prior experiments of similar amylose fragments.\textsuperscript{28} The dodecasaccharide 1 underwent numerous \( \mu \)s timescale transitions between helical and coiled shapes (Fig. 2, \textit{vide infra}), which have been hypothesized to exist based on aqueous NMR experiments.\textsuperscript{19}

In 1–5, the single major \( \alpha(1 \rightarrow 4) \) glycosidic linkage \( \psi \)n conformer was centered on \( \langle \phi, \psi \rangle \approx (70^\circ, 90^\circ) \), in agreement with the consensus from X-ray and solution NMR studies.\textsuperscript{20,29–31} Almost half of the linkages (28/62) underwent reversible \( \mu \)s exchange between this and a second minor \textit{anti} conformer centered on \( \approx (70^\circ, -60^\circ) \). These geometries agree with major and minor conformers inferred from ns molecular dynamics simulations and \textit{ab initio} calculations of \( \alpha(1 \rightarrow 4) \) linked Glc di- and tetrasaccharides.\textsuperscript{26,32} The \textit{anti} conformer, while consistent with the band-flipped geometry in cycloamyloses observed by X-ray diffraction,\textsuperscript{14,15} was predicted to be populated for at most 4\% (see ESI\textsuperscript{7}) in the 10 \( \mu \)s trajectories (with the exception of the non-reducing end linkage in 4, 9\%). Such low populations are unlikely to be detectable by standard spectroscopic experiments, perhaps explaining why they have not been reported previously. Further experimental work is deemed necessary to confirm these predictions.

The pyranose \( ^4C_1 \) chair pucker was predominant in all Glc rings from the simulations of 1–6 (see ESI\textsuperscript{7}). Pyranose puckering approached conformational equilibrium in all Glc rings after several \( \mu \)s, in agreement with recent simulations.\textsuperscript{17–19,27,33,34} The majority of pyranoses (63/70) reversibly transitioned from the most populated \( ^4C_1 \) chair to the inverted \( ^1C_4 \) pucker. Rings populated non-\( ^4C_1 \) puckers for 1–15\% during the simulations, consistent with analysis of 174 Glc-protein X-ray co-complexes.
(<2 Å resolution), where 11% contained non-\( ^{\text{2}}\)C\(_{1}\) Glc shapes (see ESI†). Resonance overlap and conformational averaging compound precise experimental quantification of such low pyranose pucker populations by NMR. Predicted ring \(^{1}\)H-\(^{1}\)H spin-vicinal couplings (\(J_{\text{HH}}\)), however, were in acceptable agreement with prior measurements for free 1-O-methyl-Glc\(^{35}\) and a maltose disaccharide,\(^{36}\) which are not indicative of perfect \(^{4}\)C\(_{1}\) Glc chairs (see ESI†). The trend for computed ring \(J_{\text{HH}}\) values to be lower than experiment (and to deviate by up to 2 Hz) was consistent with previous comparisons using GLYCAM06.\(^{17,34}\) This may be attributable to slight differences in chemical environments (simulation vs. experiment) or force field inaccuracies.\(^{17,34}\)

These analyses show agreement between the \(\mu\)s simulations of 1–6, X-ray, solution NMR, ns molecular dynamics and \textit{ab initio} computational studies. Experimentally observed shape, glycosidic linkage and pyranose ring conformers were sampled extensively while hitherto unquantified \(\mu\)s conformational exchange was predicted. The simulations lead us to conclude that modeling of \(|1\rightarrow 4|\) linked Glc oligosaccharides on \(\mu\)s timescales is both essential and valuable, enabling quantitative predictions of currently enigmatic helix-coil, glycosidic linkage and pyranose ring exchange kinetics.

**Polymer kinetics in a dodecasaccharide model**

In 1, glycosidic linkage \(\text{syn} \leftrightarrow \text{anti}\) exchange occurred more frequently towards the middle of the dodecasaccharide. Computed forward rates (Fig. 3) ranged from \(\approx 10 \text{ s}^{-1}\) (at the termini) to \(\approx 70 \text{ s}^{-1}\) (at the chain center). Over 10 \(\mu\)s, only two such transitions were present in the hexasaccharide 4 (in the central and non-reducing end linkages) and just one occurred in the trisaccharide 5 (at the non-reducing end). The simulation of 5 was extended to 20 \(\mu\)s and a single further glycosidic linkage \(\text{syn} \leftrightarrow \text{anti}\) transition occurred (see ESI†).

The terminal Glc rings of 1 underwent forward (\(^{4}\)C\(_{1}\) \(\rightarrow \) \(^{4}\)C\(_{1}\)) chair–chair exchange at a rate of \(\approx 1–2 \text{ s}^{-1}\). All internal rings inverted more frequently, in the range \(\approx 4-5 \text{ s}^{-1}\) (Fig. 3). This end-effect was not apparent in the shorter hexa- and tri-saccharide (4 and 5, respectively) where forward ring transitions were within \(\pm 0.3 \text{ s}^{-1}\) of each other and chair–chair exchange occurred at notably slower rates compared to in 1 (\(< 1 \text{ s}^{-1}\)). The single exception to this trend was the pyranose adjacent to the reducing end in 4, which inverted at a rate of 2 \(\mu\)s\(^{-1}\). The monosaccharide ring 6 inverted at a rate of 0.9 \(\mu\)s\(^{-1}\).

The two dodecasaccharide terminal pyranoses had the inverted \(^{1}\)C\(_{4}\) chair as the second most favored pucker, with computed relative free energies (\(\Delta G^0, \text{cal/mol}\)) of 1.7 and 2.3 kcal mol\(^{-1}\) at the reducing and non-reducing ends, respectively. At the termini, the skew-boat \(^{0}\)S\(_{2}\) was the third most populated Glc ring conformation. However, at all internal residues \(^{0}\)S\(_{2}\) was preferred over \(^{1}\)C\(_{4}\) and the free energy separating the puckers was greater at \(\approx 3–4 \text{ kcal mol}^{-1}\). Compared with the two terminal pyranoses, the internal Glc rings explored a greater number of chair puckers (see ESI†).

In 1–6, the rates of \(\mu\)s ring inversions and linkage \(\text{syn} \leftrightarrow \text{anti}\) exchange (1–5 only) were affected by the chain position and the degree of polymerization (as seen previously in \(\mu\)s simulations of human oligosaccharides\(^{17,19,27,33,34}\)). Forward pyranose chair–chair exchange rates were less dependent on chain location and length than the linkage \(\text{syn} \leftrightarrow \text{anti}\) kinetics. Comparison of puckering in 6 (free \(\alpha\)-anomer) with a \(\mu\)s study of the \(\beta\)-D-glucose monosaccharide\(^{35}\) (which underwent forward transitions at a rate of 2 \(\mu\)s\(^{-1}\)) implicates anomic configuration in \(\nu\)glucose ring flexibility (as for idose\(^{38}\)).

The increased rates of internal (cf. terminal) linkage and ring transitions in 1 (Fig. 3), and also the greater flexibility at the chain center in longer fragments (comparing 1, 4 and 5), suggests that the augmented kinetics result from increased mass, concomitant additional momentum and force acting on the chain center. The small range of internal Glc forward inversion rates, across the tri-\(< 1 \text{ s}^{-1}\), hexa-\(< 1–2 \text{ s}^{-1}\) and dodecasaccharides \(\approx 4–5 \text{ s}^{-1}\), suggests that chair–chair transitions may have reached a maximum rate in 1. Thus the dodecasaccharide ring kinetics are likely to be representative of polymeric puckering behavior. The large difference in internal linkage \(\text{syn} \leftrightarrow \text{anti}\) transition rates, comparing the hexasaccharide 5 (one transition in 10 \(\mu\)s) with the dodecasaccharide 1 (\(\approx 70 \text{ s}^{-1}\)), suggests that even faster rates for this exchange could manifest in longer chains. Despite this possibility, the 10 \(\mu\)s simulation of 1 represents the most realistic set of atomic kinetic predictions for single-stranded amyllose polymers to date.

**Microscopic flexibility underpins helix-coil exchange**

The predominant conformers from the simulation of 1 were helix-like (58% of the simulation) with computed \(R_g\) values in the range 12–14 Å (Fig. 2A–C). The time series for \(R_g\) (Fig. 4A) revealed \(\mu\)s exchange between extended shapes and meta-stable self-associating coiled conformers with \(R_g \approx 8 \text{ Å}\) (e.g., Fig. 2D and E). These most compact structures persisted for up to \(\approx 0.5 \mu\)s and were associated with a loss of water from the first hydration shell (see ESI†). Transient intermediate conformers contained loop-like and semi-helical sections (e.g., Fig. 2F–J). Helix-coil exchange occurred at forward (helix \(\rightarrow\) coil) and

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**Fig. 3** Forward aqueous transition rates for (A) glycosidic linkages (\(\psi, \text{syn} \leftrightarrow \text{anti}\)) and (B) pyranose rings (\(\phi, \text{anti} \rightarrow \text{syn}\)) in a model amylose dodecasaccharide 1. Linkage exchange rates (black) were derived by counting transitions of \(\psi\) from positive (syn) to negative (anti) values. Ring inversion rates (grey) were derived by counting as \(\phi\) exchanged from \(\leq 60^\circ \) (\(^{4}\)C\(_{1}\)) to \(\geq 120^\circ \) (\(^{4}\)C\(_{1}\)). Molecular properties were calculated from a 10 \(\mu\)s unbiased explicit solvent simulation.
observation is highlighted in Fig. 5 during the initial 1 µs timeframe. Conformational exchange was increasingly prevalent towards the middle of the dodecasaccharide linkage and ring conformational exchange were correlated with glycosidic linkage 6 (numbered from the reducing end). See ESI† for time series of all kinetic insights into amylose motions.

The predominant helical nature of 1 suggests that in water single-stranded amylose is primed for encapsulation of guest molecules into the helix. While recent ns simulations predicted that V-amylose helices can collapse on short (ns) timescales,22 these molecular dynamics were incapable of exploring the µs helix-coil exchange proposed here. The relatively long-timescale transitions in \( R_g \) (Fig. 4A) concur with experimental evidence for sub-ms iodine induced helix self-assembly13 but the rate of helix-coil exchange has remained elusive, underscoring that the simulation of 1 provides a route to novel atomic scale kinetic insights into amylose motions.

The correlation between linkage syn ↔ anti exchange and \( R_g \) in 1 (Fig. 5) is as expected, based on X-ray studies that associate glycosidic linkage anti conformers with coiled cycloamylose shapes.14,15 Ring flexing in 1, in particular the presence of skew-boat puckers in coiled conformers (Fig. 5), is consistent with previous atomic force microscopy measurements that implicate non-chair ring geometries in amylose elasticity16 and also the observation of non-chair puckers in high-resolution Glc-protein co-complexes (vide supra).

The simulation of 1 suggests that the microscopic basis of helix-coil exchange, which underlies gelation and the partial crystallinity of starch granules, involves correlated µs conformational transitions in both linkages and rings. These exchange rates have hitherto been inaccessible to precise quantification by experiments or ns timescale simulations. Based on these computationally-derived insights, it is concluded that GPU-accelerated molecular dynamics simulations are a useful theoretical procedure for assessing the kinetic properties and helical stability of derivatized amylases, and also for future studies of amylose inclusion complexes (e.g., encapsulated drugs or flavor ingredients).

**Microsecond stability of model double-helices**

The simulations of 2 and 3 contained µs fluctuations in \( R_g \) that were not repeatedly sampled (Fig. 6). The parallel double-helix 3
Initially disassembled, it reformed over \( \approx 0-1 \mu s \) with no similar event being observed in the subsequent 9 \( \mu s \). Although this double-helical collapse may be attributed to a suboptimal starting geometric configuration (illustrating that similar ns simulations would be sensitive to the initial coordinates), it was concluded that 2 and 3 had not reached conformational equilibrium over 10 \( \mu s \). While even longer simulations will be needed to confirm this, the dynamics of 2 and 3 were interpreted excluding the initial 1 \( \mu s \) to ensure that properties were derived only from shapes with predominantly double-helical tertiary structures and to facilitate direct comparisons with the model single-stranded amylose 1.

In the simulation of 2, iterative dodecasaccharide dissociation and association at the termini caused frequent reversible transitions from the predominant antiparallel double-helix (\( R_g \approx 13 \text{ Å}, \text{Fig. 7A} \)) to more contracted partially unwound shapes (\( R_g < 12 \text{ Å}, \text{Fig. 7E–J} \)). These conformers were short-lived (tens of ns) and the two chains did not completely dissociate during the 10 \( \mu s \) simulation of 2. The parallel double-helix 3 was comparatively stable for the majority of the 10 \( \mu s \) simulation (93\%, \text{Fig. 8A}). Its initial collapse resulted in shapes containing coiled and semi-helical dodecasaccharides (\text{Fig. 8B–J}). In light of conformers exhibiting two essentially coiled amylose chains in 3 (e.g., \text{Fig. 8F and H}), it is noteworthy that this disassembly did not result in dimerization to the antiparallel configuration.

Considered together, the simulations of 2 and 3 suggest that the parallel configuration is a more stable double-helix in water (cf. antiparallel). This finding supports the consensus view from previous X-ray studies of A type amylose\(^{37,38}\) while remaining consistent with crystallographic evidence for antiparallel double-helices.\(^{30}\)

**Tertiary organization slows linkage and ring kinetics**

As for 1, the simulations of the antiparallel and parallel double-helices 2 and 3 contained correlated \( \mu s \) conformational transitions in \( R_g \), glycosidic linkages and pyranose rings (see ESI\(^+\)). In both double-helices, the rates of forward linkage (\( \text{sym} \rightarrow \text{anti} \)) and ring (\( ^1C_1 \rightarrow ^1C_4 \)) transitions (derived excluding the initial 1 \( \mu s \)) were fastest at the dodecasaccharide termini and prevalent at the reducing ends (\text{Fig. 9}). Here, these transitions were more frequent in the antiparallel double-helix 2 (cf. parallel 3). All Glc pyranoses in 3 underwent chair–chair exchange, while ring inversion was comparatively suppressed in the central Glc rings of 2.

The finding that glycosidic linkages and pyranose rings exchanged most rapidly at the termini of 2 and 3 contrasts...
with the model single-stranded amylose dodecasaccharide 1, wherein computed forward transition rates for these degrees of freedom were faster towards the center of the chain (Fig. 3). Furthermore, linkage and ring transitions in 2 and 3 were all much slower than in 1 (comparing the computed rates in Fig. 3 and 9). Recently it was hypothesized that carbohydrate secondary-structure (pyranose stacking) restricts mobility and 9). Recently it was hypothesized that carbohydrate secondary-structure (pyranose stacking) restricts mobility and...
the gt conformer. The double-helices 2 and 3 were built manually using copies of 1, avoiding steric clashes while ensuring proximity of inter-strand hydroxymethyl and 2- or 3-position hydroxys. These hydrogen-bonds were of length 3-5 Å in the initial dimers and thus slightly longer than seen in X-ray data.\textsuperscript{30,37} This was not expected to impact the results in light of the extensive dynamics, wherein individual strands of the double-helices iteratively unwrapped and re-associated.

**Simulations**

Unbiased explicit solvent all-atom molecular dynamics simulations of 1–6 (Fig. 1) were performed for 10 µs each using ACEMD\textsuperscript{44} (Accelrys Ltd) and a single NVIDIA GTX TITAN graphics processing unit (5 was extended to 20 µs). Following initial conjugate-gradient energy minimization (1000 steps) the assemblies were heated from 0 to 298 K and then equilibrated in the NPT ensemble for 20 ns prior to 10 or 20 µs of NVT production dynamics. Data were recorded at 10 ps intervals. The velocity-Verlet integration algorithm and a hydrogen mass re-association scheme enabled a 4 fs time-step to be used without affecting the equilibrium distribution.\textsuperscript{45} This scheme has been validated and used to derive experimentally-consistent kinetic conformational models.\textsuperscript{46} Hydrogen atoms were constrained using M-SHAKE\textsuperscript{47} and electrostatic interactions were calculated via the PME method, with a grid spacing of less than or equal to 1.0 Å (in the X, Y and Z dimensions). Electrostatic and van der Waals interactions were truncated at 9 Å and the recommended\textsuperscript{19} scaling factor for carbohydrate 1–4 interactions (1.0) was employed.

**Molecular properties**

Properties of 1 (dodecasaccharide), 4 (hexasaccharide) and 6 (monosaccharide) were calculated using the complete 10 µs trajectories (i.e., one million datasets). For 2 and 3 (dodecasaccharide double-helices) the last 9 µs were used (see text and ESI\textsuperscript{1}) and for 5 (trisaccharide) the entire 20 µs simulation was employed. Radii of gyration were derived using a standard equation in VMD software,\textsuperscript{48} with standard deviations of the employed. Radii of gyration were derived using a standard equation in VMD software,\textsuperscript{48} with standard deviations of the

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Linkage and ring conformer populations were calculated by binning conformers and exchange rates by counting transitions between major conformers, as previously.\textsuperscript{17–20,27,33,34} Puckers were placed in bins of unequal spacing in the azimuthal angle \( \theta \), but equal spacing in the meridian angle \( \phi \) (ensuring that each bin occupied an equal area on the sphere). Pyranose ring \( \text{H}^1 - \text{H}^1 \) three-bond vicinal spin-couplings (\( ^3 J_{\text{H}1,\text{H}1} \)) were computed using the substituent-adjusted Karplus equations of Altona and Haasnoot.\textsuperscript{51} The Amber12 tool ptraj\textsuperscript{49} was used to perform hierarchical conformational clustering with a sampling frequency of every 50 frames (20 000 for a 10 µs simulation).

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**Notes and references**