

Research Highlighted on the Cover of *Journal of Physical Chemistry*

Previous highlight on this article, before the cover was selected:

Combined Analytics and Modeling Probing Polymer-Polymer Interactions Within Secondary Cell Walls leads to better structural insights

Background


- The molecular-level architecture of the plant Secondary Cell Wall (SCW) is elusive due to the amorphous and heterogeneous nature of lignocellulosic biomass.
- A combined experimental and computational approach is presented to bridge this gap.

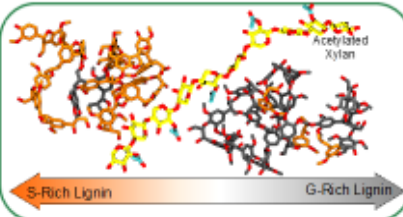
Selected Conclusions

- Polymer-Polymer interactions are probed in detail with solid-state NMR measurements (ssNMR) on ^{13}C -enriched poplar stems.
- Extent of the lignin/hemicellulose interaction is **quantified** for the first time: $\sim 70 - 75\%$ of all acetylated xylan share surface contact with lignin.
- Evidence of preferential xylan/lignin binding: S-lignin > G-lignin.
- Xylan AC_{Me} /Lignin inter-polymer distance is in the order of $0.4 - 0.5 \text{ nm}$
- Molecular dynamics simulations can illuminate the underlying atomistic architectures dictating specific ss-NMR signatures.

Significance

- Investigating plant SCW superstructure using a combined ssNMR and MD approach will greatly improve our molecular-level understanding of cell wall recalcitrance. Experimental data (ssNMR) will inform detailed atomistic cell wall models (MD), together providing better tools capable of predicting deconstruction performance and biomass structure.



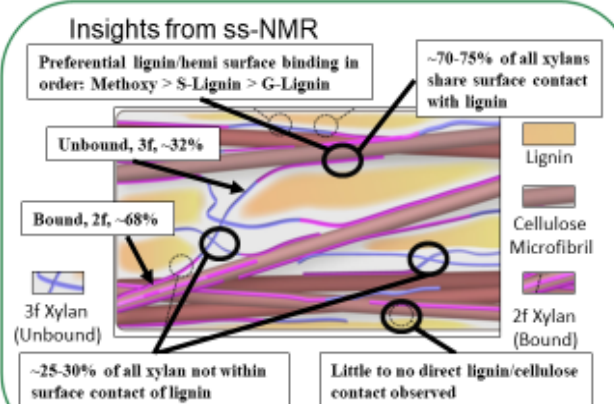


Impact of tailored xylan and lignin on ss-NMR observables from MD simulations

Insights from ss-NMR

Preferential lignin/hemi surface binding in order: Methoxy > S-Lignin > G-Lignin

$\sim 70-75\%$ of all xyans share surface contact with lignin



Unbound, 3f, $\sim 32\%$


Bound, 2f, $\sim 68\%$

$\sim 25-30\%$ of all xylan not within surface contact of lignin

Little to no direct lignin/cellulose contact observed

Adapted from: Kang and Wang, *Nat. Comm.* (2019)

B. Addison, D. Stengel, V. Bharadwaj R. Happs, C. Doeppke, G. P. Holland, T. Wang, Y. J. Bomble, A. E. Harman-Ware. 2020. *J Phys Chem B*, DOI:10.1021/acs.jpcc.0c07759 (cover article)



Informed Molecular Models of the Plant Cell Wall

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