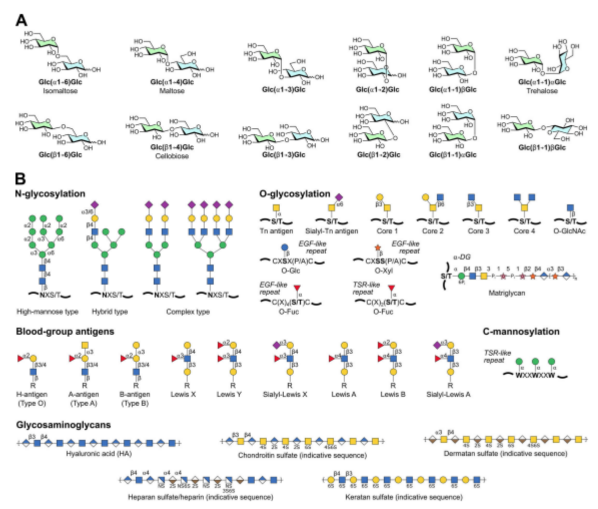


Structural glycobiology â?? from enzymes to organelles

Description

Biological carbohydrate polymers represent some of the most complex molecules in life, enabling their participation in a huge range of physiological functions. The complexity of biological carbohydrates arises from an extensive enzymatic repertoire involved in their construction, deconstruction and modification. Over the past decades, structural studies of carbohydrate processing enzymes have driven major insights into their mechanisms, supporting associated applications across medicine and biotechnology. Despite these successes, our understanding of how multienzyme networks function to create complex polysaccharides is still limited.



Molecular diversity of biological carbohydrates. (A) Regio- and stereo-isomeric combinations for a theoretical dimer of D-glucose. Eleven unique permutations are possible [Glc(α1â??1)βGlc and Glc(β1â??1)αGlc are equivalent], far exceeding the complexity of nucleic acids and proteins. (B) Selected glycan types found in eukaryotes, depicted in symbol nomenclature for glycans (SNFG) format [14]. Note that glycosaminoglycans are highly heterogenous and variable polymers â?? indicative sequences shown may not fully reflect natural complexity.

Emerging techniques such as super-resolution microscopy and cryo-electron tomography are now enabling the investigation of native biological systems at near molecular resolutions. Here, we review insights from classical in vitro studies of carbohydrate processing, alongside recent in situ studies of glycosylation-related processes. While considerable technical challenges remain, the integration of molecular mechanisms with true biological context promises to transform our understanding of carbohydrate regulation, shining light upon the processes driving functional complexity in these essential biomolecules.

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