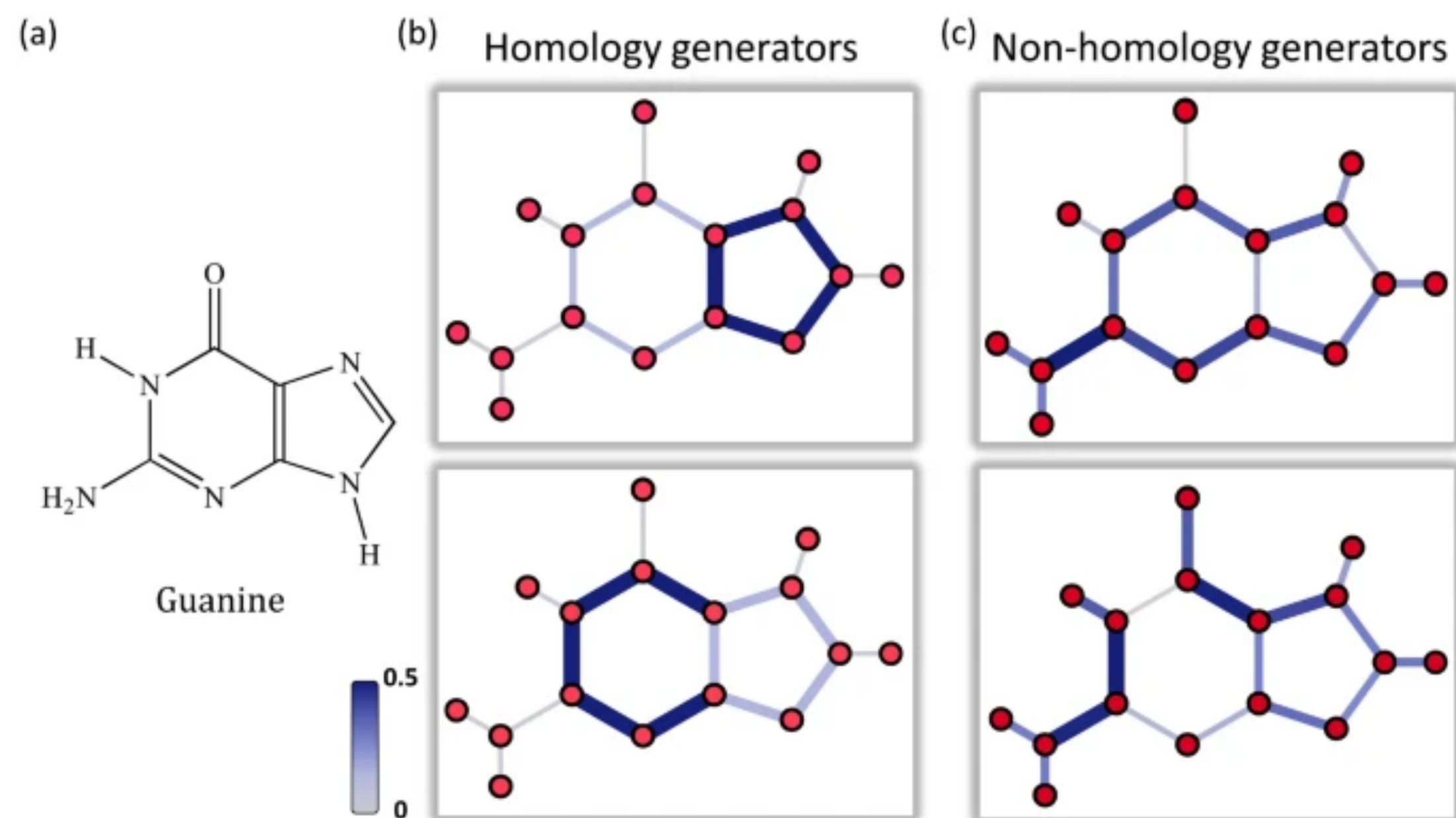
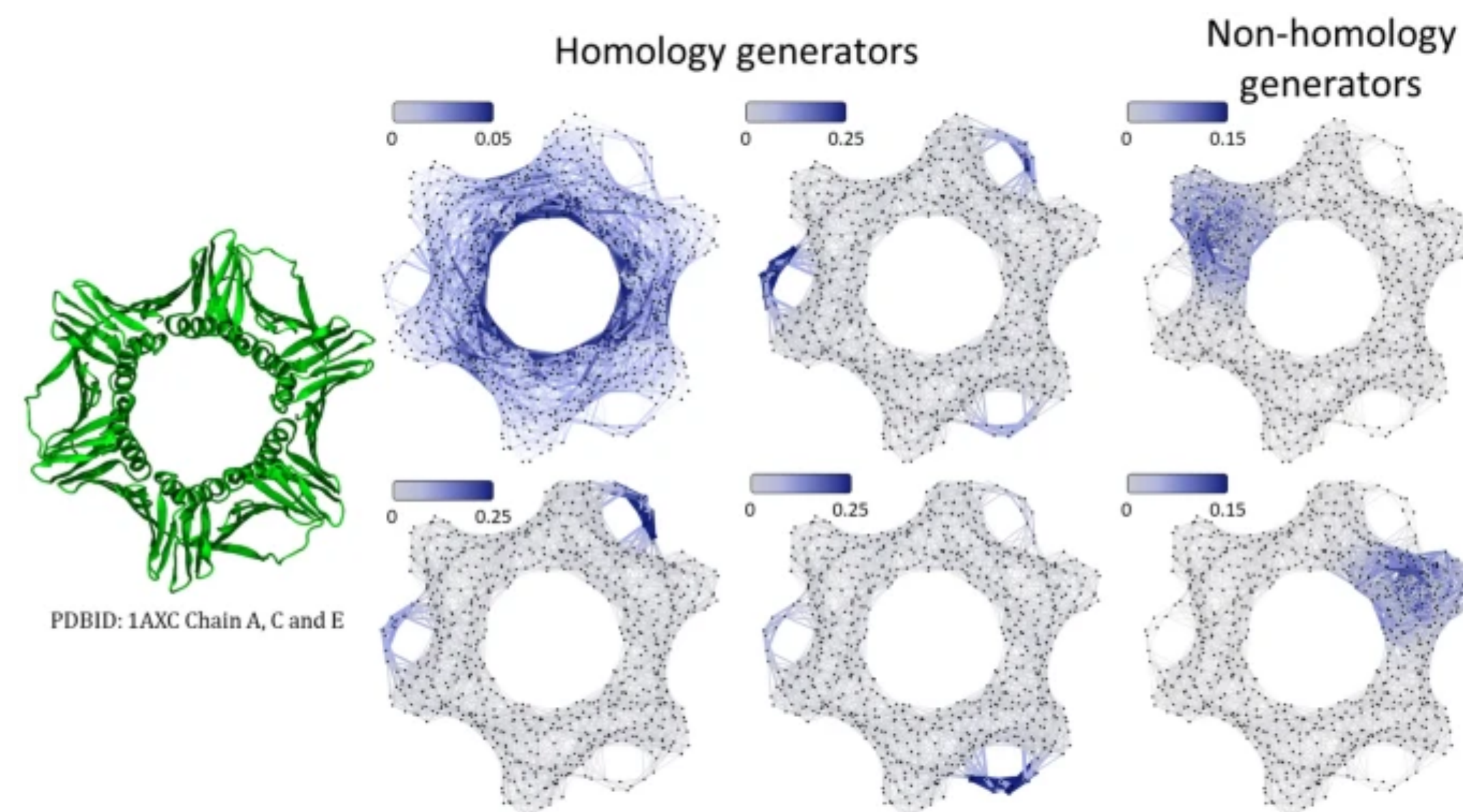


# Ideas on molecular dynamics

Wei, R.K.J., Wee, J., Laurent, V.E. *et al.* Hodge theory-based biomolecular data analysis. *Sci Rep* (2022)



(a) The graph representation for Guanine. (b) The zero-eigenvalue-related eigenvectors are homology generators. Geometrically, their largest absolute values indicate the associated loop structures. (c) Nonzero-eigenvalue-related eigenvectors are more related to domain, cluster and community structures.

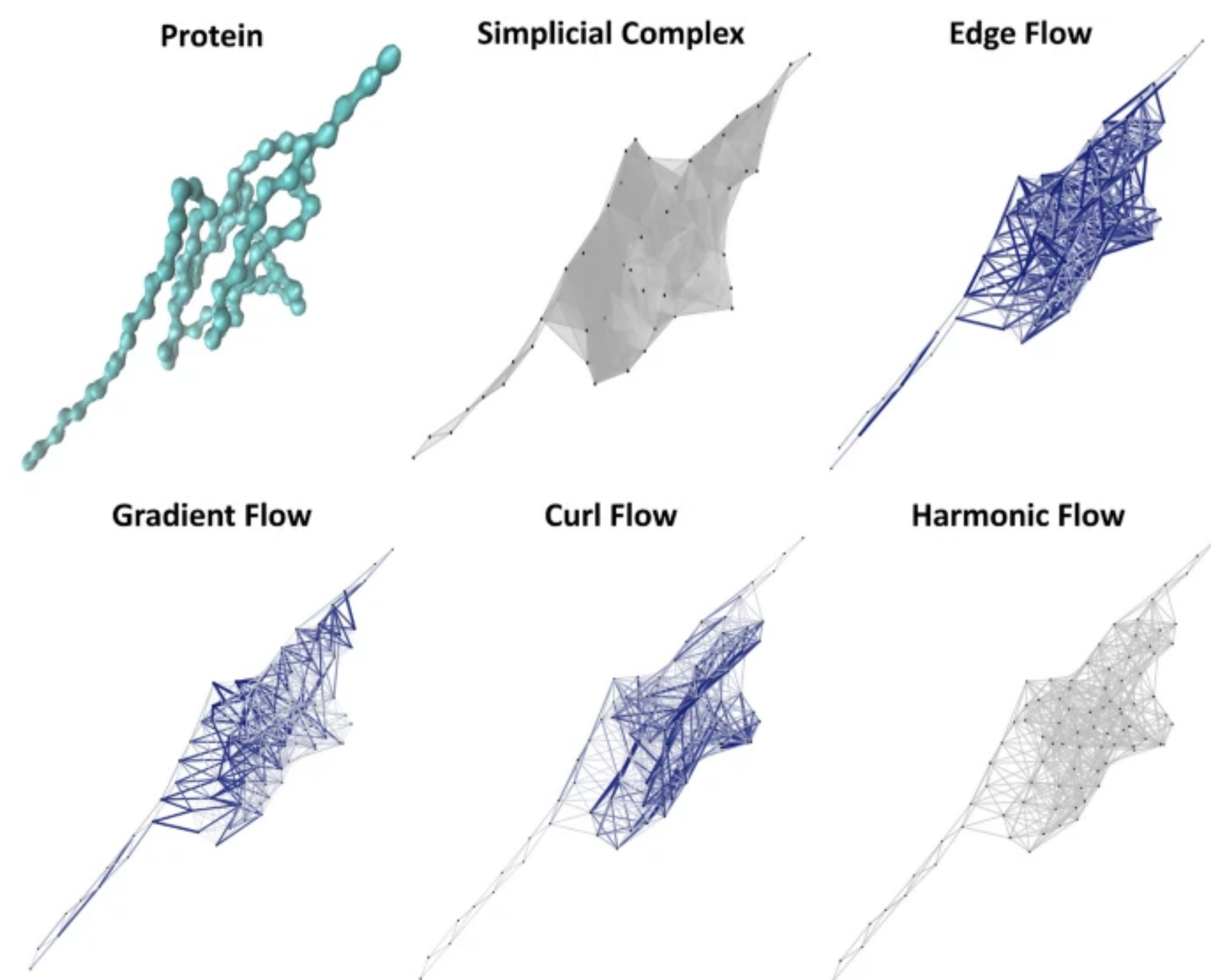


protein structure analysis for protein (ID:1AXC). Four 1D homology generators are from four zero eigenvalue related eigenvectors of 1D HL matrix. Two non homology generators are from two smallest nonzero eigenvalue related eigenvectors. Homology generators characterize loop and cycle structures, while non-homology generators indicate information about domains and communities.

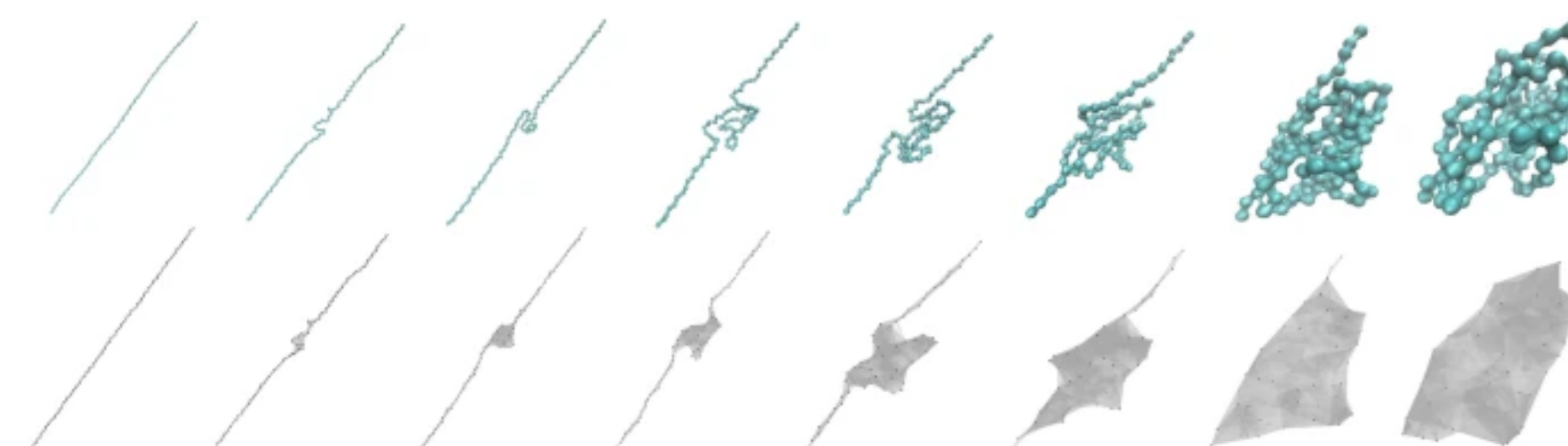


# Ideas on molecular dynamics

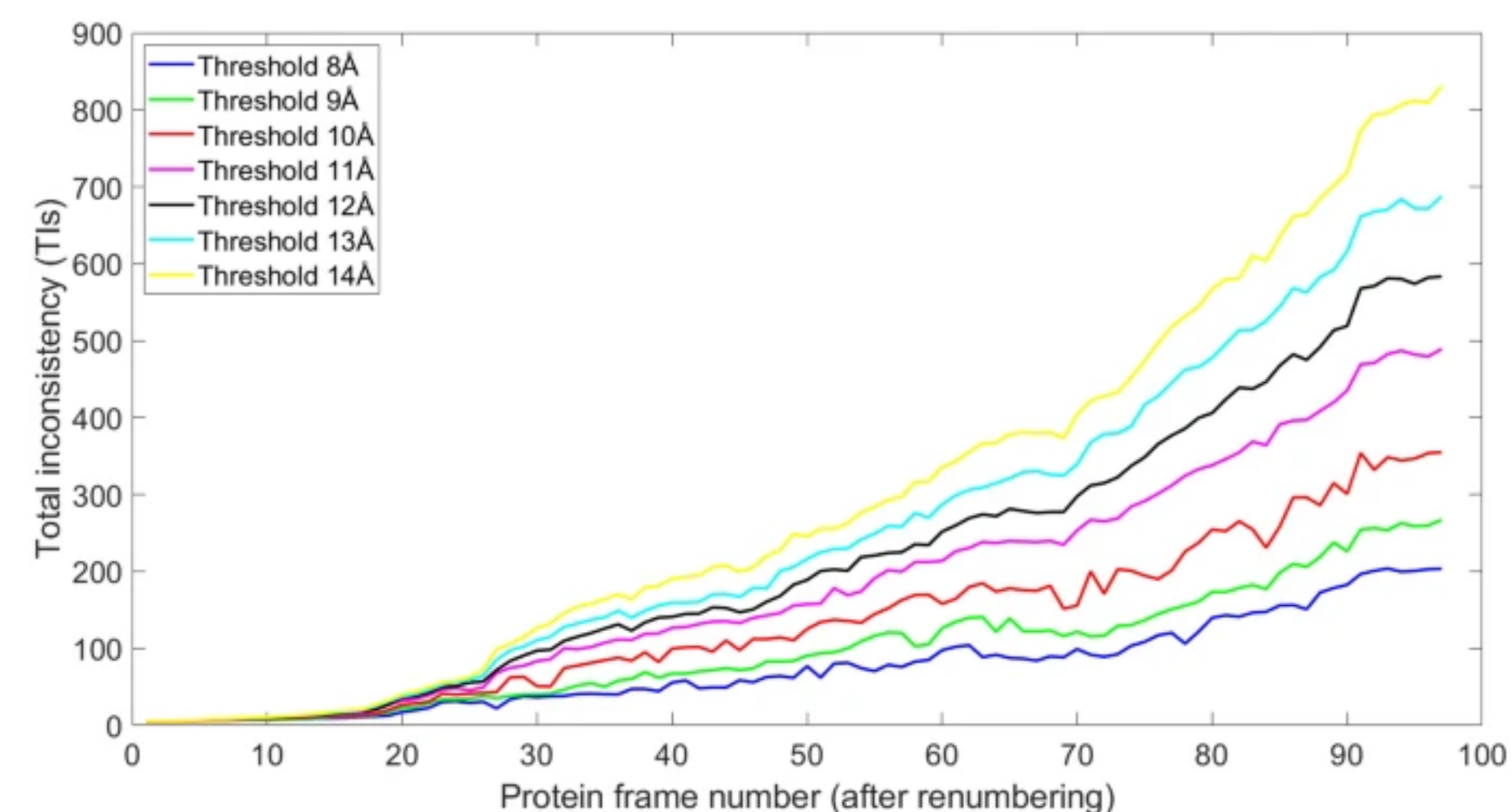
Wei, R.K.J., Wee, J., Laurent, V.E. *et al.* Hodge theory-based biomolecular data analysis. *Sci Rep* (2022)



gradient, curl, and harmonic, for a partially-folded protein



Eight configurations (after renumbering) extracted from the steered molecular dynamics simulation of Titin molecule, and their associated simplicial complexes.



(TI) for Titin configurations during the SMD simulations