SAKe: Computationally Designed Modular Protein Building Blocks for Macromolecular Assemblies

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Advances in computational protein design have allowed for the development of new proteins with unique properties. Symmetric designer proteins have remarkable stability and can serve as versatile building blocks for the creation of macromolecular assemblies. Here we present the development and structural determination of SAKe: A new symmetric, stable protein building block with modifiable loops. Following the observation of pH induced 3D self-assembly, we engineered metal binding sites along the protein’s internal rotational axis to fabricate 2D surface arrays. Using atomic force microscopy, we demonstrated Cu(II) dependent on-surface 2D self-assembly. Additionally, using dynamic light scattering and x-ray diffraction, we identified and characterized a SAKe mutant which shows in solution Zn(II) mediated nanocage formation. This work showcases a stable and highly modifiable SAKe protein scaffold, which holds promise as a building block for the creation of multi-functional macromolecular materials.

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