# Laboratory of Biomolecular Modeling and Dynamics Institute for Protein Research



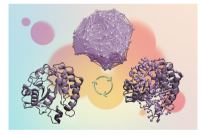
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#### Unlocking the Intricacies of Protein Structures: A Journey into Biologically Relevant Dynamics through Computational Modeling and Simulations

Proteins, the building blocks of life, are not static entities; they possess a dynamic nature, akin to any other object on Earth. At our laboratory, we delve into the fascinating world of proteins, employing structural data gleaned from advanced experimental techniques like X-ray Crystallography, cryo-electron microscopy, and small-angle X-ray scattering. Our mission? To model the myriad conformations these proteins can adopt in solution and, employing the principles of physics, simulate their flexible movements — their dynamics — deciphering the very essence of their functioning.



### Our approach is multifaceted, addressing the unique challenges posed by protein structures:

 \*\*Completing the Puzzle:\*\* Protein structures often present with missing information, requiring us to analyze analogous structures to fill the gaps.

2) \*\*Taming the Data Deluge:\*\* Copious amounts of data can overwhelm, necessitating innovative strategies. We employ techniques such as simplifying their representation through coarse-graining and utilizing methods like elastic network model-based normal mode analysis, which do not demand exhaustive details of their motions. 3) \*\*Integrating Insights:\*\* Many experimental methods provide dynamic glimpses but lack molecular resolution. We develop methods that amalgamate information from diverse sources, building a reliable understanding of the system under scrutiny.

# Our research spans a wide spectrum of intriguing topics:

- \*\*Protein Allostery:\*\* Unraveling the mysteries of protein regulation.
- \*\*Oligomerization and Biological Complex Assembly:\*\* Investigating the intricate dance of protein interactions.
- \*\*Evolutionary Conservation of Structure and Dynamics:\*\* Exploring the ancient echoes within protein architectures.
- \*\*Proteins of Interest:\*\* Delving deep into proteins like TIM Barrel fold proteins, PyrR, dihydrofolate reductase, leukotriene alpha-4 hydrolase, and more.

## Our arsenal of methods includes, but is not confined to:

 - \*\*Python Programming:\*\* Harnessing the power of coding for intricate analyses.
 - \*\*Statistical Analysis:\*\* Extracting

 \*\*Bioinformatics Tools:\*\* Extracting meaningful insights from complex datasets.
 \*\*Bioinformatics Tools:\*\* Employing advanced tools for sequence alignment and structure prediction.

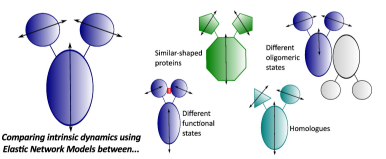
 - \*\*Molecular Dynamics Simulations:\*\*
 Simulating the dynamic dance of proteins at the molecular level.

- \*\*Normal Mode Analysis:\*\* Probing the fundamental modes of protein motion.

- \*\*Deep Learning Approaches:\*\* Leveraging the potential of artificial intelligence for nuanced understanding.

- \*\*Biomolecular Visualization and Animation:\*\* Crafting visual narratives that illuminate the intricate world of proteins.

- \*\*Web-Based Tool Development:\*\* Creating user-friendly interfaces for the broader scientific community.



Join us on this expedition, where we unravel the enigmatic world of proteins, one dynamic movement at a time. Institute for Protein Research, Osaka University 3-2 Yamadaoka, Suita, Osaka 565-0871, Japan

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Scan here for the lab's website >>

