



Short-Talk Session Abstract

<A01_Structural biology>

Fri. 1st December / 9:00 -9:40am JST / Event Hall

Light energy conversion mechanism of photosensitive membrane proteins revealed by cryo-electron microscopy

Masahiro Fukuda (The University of Tokyo)

In nature, a variety of light-sensitive proteins exist and convert light energy into various types of work. Channelrhodopsins (ChR) are representative photosensitive membrane proteins that can convert light energy into ion transport through cell membranes. Although ChRs are widely used as an optogenetic tool in fields such as neuroscience, there are still many unknown points on the molecular mechanism of ChRs. In this talk, the molecular mechanism of recently discovered unique ChRs will be presented.

Synthesis of photocaged L-tyrosine containing proteins through an Escherichia coli cell-free protein synthesis system for time-resolved structural analysis

Toshiaki Hosaka (RIKEN Center for Biosystems Dynamics Research)

Genetically encoded caged amino acids can be used to control the dynamics of protein activities and cellular localization in response to external cues. In this study, we have established a protein preparation system that introduces photocaged Tyr at arbitrary positions using an Escherichia coli cell-free protein synthesis system. Using this system, we prepared lysozyme and homoisocitrate dehydrogenase with photocaged Tyr, and crystallographic study showed that the site-specifically incorporated photocaged Tyr were degraded to tyrosine by light irradiation on the crystals. Thus, this cell-free protein synthesis of proteins with the photocaged Tyr could facilitate time-resolved structural analysis of proteins, including medically important membrane proteins.



Crystal structure of V2HeR3, a Viral heliorhodopsin transporting proton

Ritsu Mizutori (Nagoya Institute of Technology)

Heliorhodopsins (HeRs), a recently discovered rhodopsin family, are widely present in archaea, bacteria, unicellular eukaryotes, and giant viruses, but their function remained unknown. In 2022, we reported that a viral HeR from *Emiliana huxleyi* virus 202 (V2HeR3) is a light-activated proton transporter. Comprehensive mutagenesis study using electrophysiological experiment revealed some amino acids together with the retinal Schiff base forming the proton transporting gate. Here, we show the 2.1 Å resolution structure of V2HeR3. Structural and spectroscopic analyses reveal V2HeR3 specific water-mediated hydrogen bond network in both extracellular and cytoplasmic sides. These structural elements are believed to enable proton selective transport functions.

Identifying functional hotspot residues for activation in M2 muscarinic receptor (M2R)

Kota Katayama (Nagoya Institute of Technology)

The M2 muscarinic receptor (M2R) is a prototypical G-protein-coupled receptor (GPCR) that serves as a model system for understanding both ligand recognition and activation of GPCR. Here, we identify the mechanisms governing M2R activation flexibility using vibrational spectroscopy in combination with mutagenesis and organic synthetic chemistry, focusing on the native agonist acetylcholine. From these studies, we found that the exquisite distance between acetylcholine and Asn404, one of the amino acids that constitute the ligand binding site, is important for M2R activation, and that the N404Q mutant exhibits partial activity-like conformational changes. Thus, these results strongly indicate that Asn404 is a hotspot residue in M2R activation.



<A01_Chemical biology>

Fri. 1st December / 10:00 -10:40am JST / Event Hall

**Next Generation Biosensors Enabled by High-speed Visualization of
Dynamic Mechanisms**

Takuya Terai (The University of Tokyo)

Fluorescent protein (FP)-based biosensors are powerful tools in biology research, but only a few biosensors have excellent performance. A major limitation for their development is the lack of understanding of the mechanisms by which they operate. To address this issue, we are trying to capture a Molecular Movie of a crystallized FP biosensor that responds to a ligand released by light illumination. We have synthesized photocaged lactate and validated its performance. We have got grown crystals of lactate biosensors in their apo states, though the quality may need further improvement. The latest progress will be described in the presentation.

**A light-gated K⁺ channel generated by introduction of a photo-isomerizing
unnatural amino acid**

Takushi Shimomura (National Institute for Physiological Sciences)

The introduction of genetically encoded unnatural amino acids (UAAs) allows us to confer a variety of functions on target proteins. We introduced phenylalanine-azobenzene (Pab), a photo-isomerizing UAA, into KcsA, a pH-dependent K⁺ channel. Screening of the Pab position for introduction resulted in a mutant that can switch its activity with high efficiency by ultraviolet and visible light. Protein purification of the Pab mutant showed a quality similar to that of wild-type. Detailed analysis of the KcsA Pab mutant reveals the relationship between pH and light dependence. These results suggest that Pab introduction may be a useful method to confer light dependence to various proteins.



Development of adenosine A2A receptor-selective photo-switchable ligands

Tomohiro Doura (Nagoya University)

Subtype-selective photo-switchable ligands are considered a kind of powerful molecular tools for photopharmacology or optochemogenetics of G-protein coupled receptors (GPCRs). Here we report two kinds of adenosine A2A receptor (A2AR)-selective ligands with photo-switching properties. One A2AR-selective ligand is termed photoAd(vio) which is isomerized to the cis-isomer by ultraviolet (UV) light irradiation. Because the cis-isomer of photoAd(vio) activates A2AR selectively, photoAd(vio) allows to control A2AR in living cells with high spatiotemporal resolution. The other A2AR-selective ligand is named photoAd(blue). The cis-isomer of photoAd(blue) is isomerized to the trans-isomer which activates A2AR selectively by blue light irradiation. The thermally stable trans-isomer of photoAd(blue) is expected to be exploited in the structural analysis of A2AR-selective photo-switchable ligands.

Crystal structure of photoresponsive ligand bound adenosine A2A receptor

Tsuyoshi Araya (Kyoto University)

The rational synthetic expansion of photoresponsive ligands is important for the development of G protein-coupled receptors (GPCRs) photopharmacology. In this study, we synthesized the novel photoresponsive and receptor selective ligand photoNECA(blue) for adenosine A2A receptor and determined the crystal structure of the ligand-bound receptor complex. This complex structure could explain the binding mode and the photoresponsive mechanism of photoNECA(blue). And we mentioned the possibility that the photoresponsive ligand could be applied not only to photopharmacology but also to the dynamic structural analysis of GPCRs from this study.

**<A01_Structural biology>**Fri. 1st December / 9:00 -10:40am JST / Room 403**Real-Time Structural Changes during the Light-induced Water Oxidation of
Photosystem II Caught by Time-Resolved Crystallography**

Michihiro Suga (Okayama University)

Photosystem II catalyzes water oxidation by capturing sunlight energy at the unique pair of Chlorophylls called P680 within a picosecond time scale that initiates a serial reduction/oxidation reaction between two plastoquinone, QA, and QB, a radical active tyrosine residue Yz, a catalytic manganese cluster, and substrate water molecules. This reaction is the so-called S₁-state cycle catalyzed by the Mn₄CaO₅ cluster, which incorporates an extra oxygen O₆ at the S₃ state to form a possible di-oxygen. The structural changes of the metal cluster and its environment have been examined at the millisecond time range.

**An Attempt at Elucidating the Reaction Mechanism of Cytochrome c
Oxidase Using Simultaneous Measurement of XES and XRD**

Atsuhiko Shimada (Gifu University)

Cytochrome c oxidase (CcO) has four essential redox metal sites crucial for its proton-pumping and O₂ reduction reaction. It has been suggested that unidirectional proton-pumping across a membrane utilizes the structural changes of protein moieties driven by the redox changes of metal sites during the O₂ reduction reaction. To elucidate the reaction mechanism of CcO, it is necessary to comprehend the correlation between the electron states of metal sites and the protein structure. For the sake of this objective, I endeavored to simultaneously acquire XRD data (atomic coordinates) and XES data (electron states of metal sites) using XFEL.

Photoresponse mechanism of a bifunctional cryptochrome

Junpei Yamamoto (Osaka University)



Photolyase/cryptochrome superfamily (PCSf) proteins are flavoproteins bearing flavin adenine dinucleotide (FAD) as a catalytic/photoreceptive center and involved in various biological functions such as DNA repair, signal transduction, entrainment of circadian rhythm, and so on. An animal-like cryptochrome from *Chlamydomonas reinhardtii* (CraCRY) is a unique protein responsible for both photolyase and cryptochrome functions. In this study, we intended to unravel dynamic behaviors of light-dependent DNA repair and FAD photoreduction via time-resolved serial femtosecond X-ray crystallography. Intermediate structures in the ns-ms region during the FAD photoreduction of CraCRY revealed molecular mechanism of unfolding of the C-terminal region.

Fast time-resolved structural analysis of light-response in photosynthetic antenna protein Phycocyanin

Yasufumi Umena (Synchrotron Radiation Research Center Division of Synchrotron Radiation)

The cyanobacterial antenna protein, phycocyanin (PC), captures and supplies light energy to the reaction center, photosystem II. PC is composed of alpha and beta subunits of Mw 20000 with three pigment molecules, phycocyanobilin (PCB). In this study, we investigated the light-energy transfer inside the PC in sub-picosecond using pump-probe serial femtosecond crystallography (pp-SFX) with XFEL. The fast transient-absorption spectra using a PC microcrystal provided proper time-points for time-resolved pp-SFX measurement. The pp-SFX after four picoseconds showed structural changes and laser intensity-dependent damages in PCBs at 1.6 Å resolution. We will discuss the results and current issues in this study.

Reaction mechanism of photoactivated adenylate cyclase OaPAC by dynamic structural analysis

Sam-Yong Park (Yokohama City University)

Cyclic-AMP is one of the most important second messengers, regulating many crucial



cellular events in both prokaryotes and eukaryotes, and precise spatial and temporal control of cAMP levels by light shows great promise as a simple means of manipulating and studying numerous cell pathways and processes. The photoactivated adenylate cyclase (PAC) from the photosynthetic cyanobacterium *Oscillatoria acuminata* (OaPAC) is a small homodimer eminently suitable for this task, requiring only a simple flavin chromophore within a BLUF domain.

The structure of OaPAC has been capturing dynamic structural changes as molecular movies on a wide range of time scales by structural analysis using synchrotrons.

Calcium binding induced structural changes required for the photoprotein Aequorin luminescence

Toru Nakatsu (Wakayama Medical University)

Aequorin is a Ca^{2+} -binding photoprotein that was isolated from the luminous jellyfish *Aequorea victoria* and emits light by an intramolecular reaction upon binding with Ca^{2+} . Ca^{2+} binds to three EF-hand motifs of Aequorin, showing a helix-loop-helix structure. Aequorin consists of apoaequorin and 2-peroxycoelenterazine, when Ca^{2+} binds aequorin, non-covalent bounded 2-peroxycoelenterazine decomposes into coelenteramide and CO_2 to produce blue light. In this study, time-resolved X-ray crystallography of Aequorin using SACLA and SPring-8 will reveal in order to elucidate the order of calcium binding to EF-hand motifs and the mechanism of structural change associated calcium binding.

Anaerobic X-ray diffraction data collection using oxygen barrier film for study on nitric oxide reductase

Takehiko Tosha (University of Hyogo)

Bacterial nitric oxide reductase (NOR) is responsible for the elimination of cytotoxic NO. In order to elucidate the mechanism of NO decomposition at the heme/non-heme iron binuclear center of NOR, we designed experimental systems for anaerobic data collection of X-ray diffraction because the NOR-catalyzed reaction initiates the reaction of the reduced form of NOR and NO. To keep the anaerobic condition during the X-ray diffraction experiments, we focus on an oxygen barrier film to avoid the oxygen



permeation. Ethylene vinyl alcohol copolymer-based oxygen barrier film showed a potential for anaerobic X-ray diffraction experiments.

A molecular movie of photoswitchable fluorescent protein

Eiichi Mizohata (Osaka University)

The photoswitchable green fluorescent protein, rsGamillus, presents unique properties distinct from existing photo-switchable fluorescent proteins. For example, the isomeric forms of its chromophore in on/off states are trans/cis, which is opposite to that of other photo-switchable fluorescent proteins. In this study, I report the successful capture of a molecular movie of the photo-isomerization reaction for off→on switching of the rsGamillus chromophore. This was achieved using time-resolved serial femtosecond crystallography (tr-SFX) with a nanosecond laser-based pump-probe technique, capturing the events from 10 nanoseconds to the microsecond timescale.

Structure and mechanism of a Diels-Alderase and a 2-oxoglutarate-dependent dioxygenase

Shingo Nagano (Tottori University)

A Diels-Alderase, Fsa2, which is involved in equisetin biosynthesis, produces a decalin skeleton by stereo-selective 4+2 cyclization reaction. We determined the crystal structure of equisetin-bound Fsa2 and confirmed that the equisetin binding mode was consistent with the substrate binding mode previously predicted by a molecular dynamics simulation.

TauD, which hydroxylates taurine, is a non-heme iron²⁺ and 2-oxoglutarate-dependent dioxygenase. Metal-free and taurine-bound TauD microcrystals were mixed with FeSO₄ solution to induce the enzyme reaction. The occupancy of the non-heme iron was time-dependently increased, confirming the possibility of non-heme iron reconstruction of crystalline TauD. In addition, the polder omit map with a delay time of 9.0 sec confirmed the electron density extending from the carbon atom to be hydroxylated in substrate taurine toward non-heme iron, suggesting that taurine was hydroxylated in the microcrystals.



< B01_Molecular Movie Platform Design >

Fri. 1st December / 9:00 -9:20am JST / Room 404

Development of an in-vacuum diffractometer for protein micro-crystallography

Hiroaki Matsuura (RIKEN SPring-8 Center)

Microbeams available at synchrotron and FEL facilities have enabled structure determination from tiny protein crystals. However, due to fewer unit cell copies in such tiny crystals, diffraction intensities become significantly weaker. This weak signal complicates the observation of scattering from sample crystals, particularly as background scattering from the air becomes prominent. We have been developing an in-vacuum diffractometer designed for micro-crystallography with crystals smaller than 1 μ m/side. Test experiments were conducted using cytidine crystals to compare data collection under atmospheric and vacuum conditions. Our findings indicate that data collection under vacuum conditions improves crystallographic statistics such as $\langle I/\sigma I \rangle$ or R factors.

Low background graphene sample holder for XFEL-based crystallography and imaging

Akihiro Suzuki (Hokkaido University)

In the 2010s, X-ray free-electron laser (XFEL) facilities were constructed around the globe to pioneer new X-ray science by taking advantage of its highly brilliant and short radiation pulses. Single particle imaging of a protein that is difficult to crystallize was one of the driving forces behind the XFEL constructions. However, even after more than ten years in operation, detecting diffraction signals from a single particle is still a severe challenge. To improve the signal-to-noise ratio of diffraction signals, we have been developing a low-noise sample holder using graphene instead of liquid jets for sample delivery. In this presentation, we will present the current status of sample holder development and the latest result of the XFEL experiment.

<C01_Spectroscopy >Fri. 1st December / 9:20 -10:00am JST / Room 404**Stepwise calcium binding in photoluminescence protein; aequorin, revealed
by the time-resolved spectroscopy**

Tetsunari Kimura (Kobe University)

The photoluminescence protein; aequorin (AQ) has three Ca²⁺ binding sites with loop structure and the binding induces the transient emission of blue light. The stepwise Ca²⁺ binding has been proposed based on the time-resolved serial-femtosecond x-ray crystallography and serial soaking experiments. The comprehensive spectroscopic measurement is critical to evaluate the proposed mechanism. In this study, kinetic measurement of luminescence intensity along the AQ-Ca²⁺ binding and Ca²⁺ affinities were investigated by stopped-flow and isothermal titration calorimetry, respectively. Kinetic analysis clarified that the first Ca²⁺ binding induced the conformational change, which was followed by the second Ca²⁺ binding with higher affinity, indicating that the conformational change in the loop structure was necessary to accommodate Ca²⁺ ion, as proposed in structural biological experiments.

**Structural features of Lumi intermediate on a primate blue-sensitive visual
pigment revealed by FTIR**

Yosuke Mizuno (Nagoya Institute of Technology)

Cone pigments are photoreceptors responsible for color vision. Lumi, an early photointermediate, of a primate blue pigment (MB) is deprotonated at the retinal Schiff base, which is an unusual reaction not seen in other visual pigments. Here, we report the structural features of Lumi of MB measured by FTIR spectroscopy. Site-directed mutagenesis study reveals that the Schiff base proton transfers to the E113, a counterion, coupled with local perturbation of α -helices. Hydrogen bond strengths of T118 and C211 together with internal water molecules change upon the formation of Lumi, and thus these hydrogen bond alterations might be key elements enabling early proton transfer reaction.



A Unique Molecular Property of a Sodium-Pumping Rhodopsin in Krokinobacter eikastus

Yuji Furutani (Nagoya Institute of Technology)

Microbial rhodopsins exhibit various functions such as light-driven ion pumps, light sensors, light-gated channels, and light-activated enzymes. In 2013, a sodium-pumping rhodopsin was found from *Krokinobacter eikastus* (KR2). Interestingly, a substrate sodium ion is not bound near the protonated Schiff base region, which is transiently incorporated into the protein moiety upon formation of the intermediate states. Thus, the molecular mechanism has been extensively studied by using various techniques. In this short talk, I will present our recent finding regarding importance of the covalent bond connecting the retinal chromophore in KR2 (S. Ochiai et al. *Biochemistry* 2023).

Excited state dynamics in a single protein crystal by using femtosecond transient absorption microscopy

Tetsuro Katayama (Tokushima University)

Energy transfer reactions are important in photosynthetic systems as natural systems and solar cells as artificial systems. In this study, femtosecond transient absorption microscopy was utilized to elucidate the energy transfer reaction behavior between dyes in a single crystal. In this presentation, we will discuss the details of the excited-state relaxation process between dyes from the viewpoint of polarization dependence.



< C01_Computational Chemistry >

Fri. 1st December / 9:00 -10:40am JST / Room 405

Development of non-adiabatic QM/MM molecular dynamics method

Kiyoshi Yagi (RIKEN Cluster for Pioneering Research)

We have developed a non-adiabatic molecular dynamics (MD) method based on QM/MM calculations. QM/MM is a multiscale method that treats the reaction center by quantum chemical (QM) method and the biological environment as a classical force field (MM). The QM calculation of the electronic excited state, which is the computational bottleneck, is highly parallelized to achieve high performance. In addition, QM/MM is combined with a surface hopping algorithm proposed by Zhu and co-workers, which takes into account non-adiabatic transitions between electronic states. The method is applied to photochemical processes of azobenzene in solution and bacteriorhodopsin.

Overview of our research collaborations performed in the research area of "molecular movies"

Mitsuo Shoji (University of Tsukuba)

We have performed collaborative researches under the research area of "molecular movies". The target molecules and collaborators are (1) C-phycoerythrin with Y. Umena (A01), (2) copper amine oxidase with T. Murakawa (A01), (3) heliorhodopsin with K. Katayama and H. Kandori (A01), (4) lysozyme with B. Maity and T. Ueno (B01), (5) 2-oxoglutarate-dependent dioxygenase with S. Nagano (A01) and (6) P450 sesamin with E. Mizohata (A01). We overview all of these research progresses, achievements and remaining issues.

How the mGLU1 receptor dimer be activated: a molecular dynamics study

Duy Phuoc Tran (Tokyo Institute of Technology)

We present in this study our recent progress in understanding the activation mechanism of the mGLU1 receptor. In collaboration with Kiyonaka group, we have figured out the



mechanism of the activation of the mutations and ligands. In addition, we reconstructed the full-length structure of the mGLU1 dimer with Gi1 protein by molecular docking and simulation. The data show the differences in interaction interfaces between mGLU1 and the other mGLU receptors in sub-family.

Complementing molecular movie by analyzing free energy landscape, pathways, and flux of conformational change

Akio Kitao (Tokyo Institute of Technology)

We conducted molecular simulation of biomolecular systems to complement molecular movie experimentally determined. Using parallel cascade molecular dynamics (PaCS-MD) simulation and other computational techniques, we investigated conformational change of proteins by analyzing free energy landscape, transition pathways and flux. We report interfacial activation of *Candida antarctica* Lipase B and molecular mechanisms of structure support and smooth rotation in the bushing/rod complex of bacterial flagella.

Dynamical Analysis for Simulations of Orexin 2 Receptor Using Relaxation Mode Analysis

Ayori Mitsutake (Meiji University)

We have introduced and developed algorithms for protein simulation. For dynamics and stability, we introduced relaxation mode analysis and 3D-RISM theory, respectively. 3D-RISM theory estimates the solvation free energy. We use molecular dynamics simulations and these analyzes to study the inactive and active conformations of the orexin-2 receptor. From simulations of constitutively active mutants, we identified a good indicator for distinguishing between inactive and active conformations of class A GPCRs, the distance between Ile48.CA (3.46) and Tyr364.CA (7.53). From simulations of active and inactive structures in the presence and absence of ligands, we investigated characteristic amino acids that are thought to be important for activation of orexin-2 receptors.



Theoretical study on molecular mechanism of an activation process of aequorin bioluminescence

Shigehiko Hayashi (Kyoto University)

Aequorin is a bioluminescent protein which binds coelenterazine as a light emitting molecule. A chemiluminescence process of coelenterazine with a molecular oxygen in the protein binding pocket is triggered by binding of calcium ions at EF-hands of the protein distant from the binding pocket. We theoretically investigate molecular mechanism of the chemiluminescence process in the protein by means of hybrid QM/MM molecular simulations. We found significant conformational changes of the binding pocket upon the binding of calcium ions and a proton exchange through hydrogen-bond network formed by the conformational changes which produces a stable intermediate for chemiluminescence.

Unification of molecular movies and quantum molecular dynamics

Junichi Ono (Waseda University)

Proton transfers associated with structural changes play a vital role in function. Recently, molecular movies of structural changes in biomolecules have been successfully captured using time-resolved SFX at XFEL. However, direct observation of proton transfers has not been accomplished because of the limitation of spatio-temporal resolution. Here, large-scale quantum molecular dynamics (QMD) simulations were performed on the basis of the molecular movies with focusing on the chemical reactions in rhodopsins, where all the atoms were treated quantum-mechanically. In this presentation, we discuss our results demonstrating that our QMD simulations compensate for the molecular movies to reveal the microscopic mechanisms of chemical reactions.

Resi-DEM: A Comprehensive Tool for Automated Density Extraction, Identification, and Analysis of Difference Densities in Isomorphous Maps, with Occupancy-Estimation for Conformer Fitting

Sriram Srinivasa Raghavan (RIKEN Center for Computational Science)



We will introduce Resi-DEM, an new tool designed to automate the identification of difference densities in isomorphous difference density maps, further associating them with relevant residues through a clustering approach. Utilizing the clustered difference density features, we estimated the occupancy of the triggered state population via the structure factor extrapolation technique. This clustering technique was employed to organize difference densities, enabling us to represent them as networks, facilitating the tracking of density changes over various time periods. Moreover, we employed molecular dynamics trajectory sampling to determine the most fitting conformation corresponding to the triggered data set. The results were validated using metrics such as the Real Space Correlation Coefficient (RSCC) and Real Space Difference Density Z score (RSZD).

Analysis on the dynamics of bacteriorhodopsin using MD simulation

Keiko Shinoda (The Institute of Statistical Mathematics)

In the "molecular movies", we have developed force fields for lipid molecules in the archaeal membrane and constructed an archaeal model membrane. Using this, we have carried out molecular dynamics (MD) simulations of bacteriorhodopsin. In this presentation, in addition to the previously reported results on the resting state and at 16ns after excitation of bacteriorhodopsin, I will report on the results at 760 ns and 2 μ s after excitation and present a comprehensive analysis on the dynamics of bacteriorhodopsin.