Microsymposia

MS.08.2  

**Molecular basis for rotational switching in the bacterial flagellar motor**

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The bacterial flagellar motor is one of the most efficient rotary motors known to man. It rotates at hundreds of revolutions per second, yet can reverse its direction in less than one millisecond [1], [2]. Both of these attributes facilitate the rapid movement of bacteria towards favourable environments. The motor uses the potential energy from an electrochemical gradient of cations [3] across the cytoplasmic membrane to generate torque. A rapid switch from counterclockwise to clockwise rotation determines whether a bacterium runs smoothly forward or tumbles to change its trajectory [4], [5]. A protein called FliG forms a ring in the rotor of the flagellar motor that is involved in the generation of torque [6], [7] through an interaction with the cation channel forming stator subunit MotA [8]. FliG has been suggested to adopt distinct conformations that induce switching but these structural changes and the molecular mechanism of switching are unknown. We have recently determined the X-ray structure of the full-length FliG protein from *Aquifex aeolicus* [9], identified conformational changes that are involved in rotational switching and uncovered the structural basis for the formation of the FliG torque ring. This allowed us to propose a model of the complete ring (Fig. 1) and switching mechanism in which conformational changes in FliG reverse the electrostatic charges involved in torque generation.

**Figure 1:** Proposed model of the torque ring of the *A. aeolicus* flagellar motor consisting of 34 FliG proteins. Charges known to be involved in torque generation and switching are located at the outer circumference of the ring [9].


Keywords: macromolecular complex, rotary motor, bacterial locomotion

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MS.08.3  

**Structure and function of protein-based metabolic organelles in bacteria**

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Although it is not widely appreciated, many bacterial cells contain giant protein-based organelles that make it possible to carry out certain metabolic processes in a sequestered environment [1,2]. The simplest and best characterized type of bacterial microcompartment (BMC) is the carboxysome, which in cyanobacterial cells encapsulates the enzymes RuBisCO and carbonic anhydrase within a capsid-like protein shell in order to enhance the rate of CO_2_ fixation. Other more complex types of bacterial microcompartments are produced under specific conditions by well-studied bacteria such as *Salmonella* and some strains of *E. coli*. These more complex microcompartments typically encapsulate a larger set of enzymes, which metabolize a small molecule (like ethanolamine) without allowing the escape of a volatile or toxic metabolic intermediate (like acetaldelyde). The main (so-called BMC) shell proteins form hexamers, which further assemble in side-by-side fashion to form a tightly packed molecular layer. Other minor proteins form pentamers that are believed to sit at the vertices of the roughly icosahedral shell. Intact microcompartments are roughly 100 nm in diameter, with an outer shell composed of a few thousand shell proteins and an interior containing a similar numbers of enzyme molecules. Crystal structures of numerous shell proteins from various microcompartments are shedding light on principles of protein assembly and evolution, and mechanisms of molecular transport of substrates and products. Current crystallographic work will be presented.

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MS.08.4  

**Unlikely crystals: poxvirus spheroids in vivo crystallization**

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Unlikely crystals: poxvirus spheroids in vivo crystallization

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Keywords: protein assembly, symmetry, metabolism
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While protein crystallization of highly purified proteins has been studied for decades and used to painstakingly produce crystals for structural analysis, common insect viruses have evolved proteins that readily crystallize in vivo despite the complexity of the cellular environment. Viral, in vivo, crystals are among the most striking examples of protein self-assembly, ultimately leading to the formation of ultra-stable microcrystals occupying most of the infected cells.

Recently, the first structures of in vivo crystals were elucidated from viral polyhedra produced by RNA [1] and DNA [2, 3] viruses. The function of polyhedra is to package up to hundreds of viral particles constituting the main infectious form of the virus and allowing the virus to persist for years in the environment like bacterial spores.

Apart from polyhedra, spheroids produced by poxviruses are the only other known example of viral infectious crystals. RNA and DNA virus polyhedra share nearly identical lattices, similar function and, of course, common polyhedral morphologies. In contrast, whether spheroids were single crystals remained unclear because of their ovoid shape and the large size of their 100kDa matrix protein.

To elucidate the molecular organization of the third class of infectious crystals, we determined the 2.5 Å structure of entomopoxvirus spheroids. These diffraction experiments were carried out on spheroids with maximum dimensions of 15 μm directly purified from infected insects and stored frozen for over two decades.

The structure determination of spheroids confirmed that they are also single crystals despite their unusual ovoid shape. In the crystal, the main constituent of spheroids, called the spheroidin protein, adopts a multi-domain fold with five independent structural domains unrelated to either of the two types of polyhedrin protein. The asymmetric unit of the crystal contains two 100kDa spheroidin molecules adopting conformations that differ by hinge movements between the five domains. This apparent flexibility of the spheroids building blocks was totally unexpected given that crystallization occurs readily in vivo despite the complexity of the cellular milieu and results in remarkably stable crystals. A detailed analysis of the crystal organization provides an explanation to this conundrum revealing that building blocks interlock into the crystalline lattice using an extensive network of disulide bonds.

In conclusion, this structure provides a detailed view of what may constitute the most improbable protein crystals. Spheroids not only harbor an atypical ovoid shape and host hundreds of irregular virus particles without disruption of the crystalline lattice but they are also able to assemble in vivo from large and flexible building blocks. Mirroring the complexity of poxviruses themselves, spheroids are undoubtedly the most elaborate viral armors providing a unique model of in vivo crystallization.


Keywords: microcrystallography, virology, assembly

MS.09.1


Generating functions for structure and chemical composition
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Any crystal structure may be represented by a weighted chromatic digraph, the vertex set of which represents atoms and the edge set of which represents chemical bonds. We may write tetrahedrally coordinated cations and their associated anions as \( T_x \Theta_y \). For \( T_x \Theta_y \) to be a chain or ribbon, \( 5n \leq m \leq 6n \), and we may write as \( m = 5n^+ \), where \( N \) is an integer. Within the \( T_x \Theta_y \) unit, we may recognize three types of anion vertices: (1) bridging anions, \( \Theta^a \), that are bonded to two T cations; (2) apical anions, \( \Theta^b \), that are involved in linkage to other cations out of the plane of the bridging anions; and (3) linking anions, \( \Theta^c \), that link to non-T cations in the plane of the bridging anions. We can incorporate the connectivity of the cations into our algebraic representation of the chain as follows: \( (T_x \Theta^a \Theta^b \Theta^c)^n \) where \( a + b + c = 5n^+ \). The apical anions of the T-layer map onto a 6n-well which, in turn, maps onto the 3n^+ of anions of the O-layer. We may use the handshake dilemma of graph theory to examine the interaction between the two types of layers, and write a Structure-Generating

MS.08.5


X-ray structure of a functional full-length dynein motor domain
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Dyneins are microtubule-based motor complexes that power a wide variety of biological processes within eukaryotic cells, including the beating of cilia and flagella, cell division, cell migration, and the intracellular trafficking. Compared to the other cytoskeletal motors kinesin and myosin, the molecular mechanism of dynein is still poorly understood, in part due to the lack of high-resolution structural information. Because dyneins belong to the AAA+ family of mechanochemical enzymes, its structure and mechanism must be fundamentally different from the G-protein related kinesins and myosins. The X-ray crystallography of dynein or even its motor domain—a 380-kDa portion of the heavy chain responsible for dynein’s motor activity—has been challenging due to its large size and molecular complexity.

Here, we report an X-ray crystallographic analysis of the entire functional 380-kDa motor domain of Dicystostelium cytoplastymbic dynein, the longest polypeptide (~3,300 residues) that has been crystallized so far. Diffraction from crystals extends to 4.0 Å using synchrotron radiation from an undulator source (BL44XU, SPring-8). Diffraction is consistent with space group \( P2_12_12_2 \), with unit cell dimensions of \( a = 201.17 \, \text{Å}, b = 228.96 \, \text{Å}, c = 195.73 \, \text{Å} \). Based on an electron density map calculated from the TaBr_4 derivative data at 4.5 Å resolution, a helical model of the dynein motor domain has been created [1]. The analysis reveals detailed architectures of functional units responsible for the motor activity, such as the ATP-hydrolyzing, ring-like head composed of six AAA+ modules as well as the long coiled-coil microtubule-binding stalk, the force-generating rod-like linker and some unpredicted structures likely to be key to function. This long sought crystal structure provides the framework to understand a large volume of data obtained by electron microscopic, biochemical and single-molecule studies, and opens the door to detailed understanding of how dynein produces force and movement.


Keywords: biomacromolecule, biophysics, mechanism

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