Merits of the double-stranded form of the actin filament revealed by structures of the filament ends

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ACTIN FORMS A DOUBLE-STRANDED FILAMENT, AND THE MAJORITY OF ACTIN FILAMENTS IN THE CELL UNDERGO THE DYNAMIC PROCESS OF POLYMERIZATION AND DEPOLYMERIZATION AT BOTH ENDS. ACTIN DYNAMICS PLAYS NUMEROUS IMPORTANT ROLES IN EUKARYOTIC CELLS. IN ORDER TO UNDERSTAND ACTIN DYNAMICS, STRUCTURAL ELUCIDATION OF THE ACTIN FILAMENT ENDS IS PARTICULARLY IMPORTANT BECAUSE POLYMERIZATION AND DEPOLYMERIZATION OCCURS ONLY AT THE ENDS. WE HAVE DEVELOPED A METHOD TO DETERMINE THE END STRUCTURE BY CRYO-ELECTRON MICROSCOPY AND IMAGE ANALYSIS PROCEDURES. THE TWO STRUCTURES THAT WERE DETERMINED, THE ACTIN-CAPPING PROTEIN (CP) COMPLEX AND THE BARE POINTED END OF THE ACTIN FILAMENT, REVEALED UNKNOWN REGULATORY MECHANISMS OF THE ACTIN DYNAMICS. IN THESE MECHANISMS, THE ACTIN FILAMENT TAKES ADVANTAGE OF ITS OWN DOUBLE-STRANDED FORM IN THREE DIFFERENT WAYS.

1. The double-stranded form is required for end-binding proteins to recognize and bind, and these proteins do not recognize actin monomers (Fig. 1). Many end binding proteins such as CP, formin, tropomodulin and spire, regulate actin dynamics because these proteins are most effective at binding to the filament ends where the polymerization and depolymerization occur. Therefore, recognition of the end by an end binding protein is important. The actin-CP complex structure represents a simple and sophisticated manner of the recognition process (Fig. 1A–C). We believe many end binding proteins recognize the target end by a similar manner, simultaneously binding to two regions which are exposed only at the target end on two subunits located on different strands. When the filament is single-stranded, the specific recognition of the target end is more difficult (Fig. 1D–F).

2. End binding proteins can regulate the stability of the whole filament (Fig. 2) when the filament has a double-stranded form. The actin filament in muscle is very stable, whereas it is truly dynamic in the...
Figure 1. The double-stranded form is useful for end binding proteins to recognize the target end. A–C: The binding mechanism of the Capping Protein.4 (A) A three dimensional map of the actin-CP complex with fitted atomic models of CP (in red and orange) and actin molecules (in purple, blue, cyan and green). CP binds to only the barbed end, neither to the side of the filament, nor the pointed end, nor the actin monomer. (B) A schematic illustration of CP (in cyan) binding to the barbed end. The major binding site on CP, illustrated as a blue ellipse, binds to the two end subunits on the two strands simultaneously. The binding sites on the actin filament, illustrated as red ellipses, are exposed only at the barbed end, neither at the pointed end nor the side of the filament, thereby illustrating how CP recognizes the barbed end. (C) CP does not bind to actin monomers. CP requires two binding sites on two different strands with proper relative positioning for tight binding. CP can bind to only one binding site on the actin monomer even though one actin monomer has the two binding sites present (red ellipses). Consequently, the binding is significantly weaker than at the barbed end. (D–F) Schematic illustrations of a putative model on how an end binding protein (in cyan) recognizes the end when the filament is single-stranded. (D and E) Even when the end binding protein (the binding site is presented in blue) recognizes only the one end subunit (the binding site is presented in red), it can recognize the target end of the filament when the binding site on the filament is exposed only at the target end (D). However, it also binds to the monomer (E). To prevent binding to the monomer, the end binding protein (in cyan) must recognize two sites on the two subunits at the end including a site which is only exposed at the target end (F). The interaction between the end filament requires a much larger protein than with the double-stranded filament interaction model presented in (C).
lamellipodia where the turnover of the actin filament is less than one minute.\(^9\) The actin filament must cover a wide stability range and regulation of the stability by end binding proteins is essential.

(3) The double-stranded form plays an important role in defining the polarity of the dynamics (Fig. 3). One end, the pointed end, is much slower in polymerization and depolymerization than the barbed end, and these differences determine the direction of the “treadmilling” movement.\(^5\) We believe that this feature also defines the origin of the direction of the dynamics of the actin filament in a cell; where polymerization occurs at the barbed end and the depolymerization occurs at the pointed end.\(^5\)

In conclusion, our structures of the actin filament ends have revealed that the double-stranded form of the actin filament is essential in regulating actin dynamics in the cell, which is fundamental to all functions of the actin filament.

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Figure 3. The double-stranded form is important in defining the polarity of dynamics. A simplified model to explain the difference in the dynamics between the two ends is presented. A more realistic and complicated model was proposed previously in reference 5, although the concept is the same. (A) A simplified illustration of the actin monomer. The switching loop is the DNase-I binding loop in the simplified model. The actin monomer has two binding sites for the switching loop (cyan circles). One is a strong binding site (the large circle) and the other is a relatively weak binding site (the small circle). (B) A model of the barbed end. The switching loop of the end subunit (in red) binds to the strong binding site on the actin subunit in the same strand. As a result, a strong binding site for an incoming actin monomer is available and the incoming monomer can easily bind to the end. (C) A model of the pointed end. The switching loop of the adjacent subunit at the end (in red) binds to the weak binding site on the end subunit in the other strand. Consequently, the dissociation of the end subunit becomes more difficult at the pointed end because the extra binding by the switching loop (in red) prevents the end subunit from dissociating. When a new actin monomer comes in close proximity to bind to the strand, the switching loop (in red) that is already bound to a different site must first dissociate from the weaker site prior to interacting with strong binding site of the new monomer. As a result, the depolymerization and polymerization rates at the pointed end are slower than at the barbed end and represents the origin of the polarity of the dynamics of the actin filament. (D–F) When the filament is single-stranded, it is difficult to define the polarity of the dynamics. If we assume a similar switching loop in the single-stranded filament, the weak binding site for the switching loop to bind to the pointed end must be located in the same strand because there is only one strand present. However, the switching loop in the monomer can also bind to the weak binding site in the same monomer (D). In this case, the polymerization at the pointed end (E) and at the barbed end (F) is inhibited to the same extent because of the switching loop binding to the weak binding site. For the dissociation from the ends, the situation cannot be different because the connection to be severed for dissociation to occur is identical between the two ends. Therefore, the rates of polymerization and depolymerization at the both ends must be similar when the filament is single-stranded.