Overexpression of Cofilin Stimulates Bundling of Actin Filaments, Membrane Ruffling, and Cell Movement in Dictyostelium

Hiroyuki Aizawa,* Kazuo Sutoh,† and Ichiro Yahara*

*Department of Cell Biology, The Tokyo Metropolitan Institute of Medical Science, Tokyo 113, Japan; †Department of Life Sciences, Graduate School of Arts and Science, University of Tokyo, Tokyo 153, Japan

Abstract. Cofilin is a low molecular weight actin-modulating protein whose structure and function are conserved among eucaryotes. Cofilin exhibits in vitro both a monomeric actin-sequestering activity and a filamentous actin-severing activity. To investigate in vivo functions of cofilin, cofilin was overexpressed in Dictyostelium discoideum cells. An increase in the content of D. discoideum cofilin (d-cofilin) by sevenfold induced a co-overproduction of actin by threefold. In cells overexpressing d-cofilin, the amount of filamentous actin but not that of monomeric actin was increased. Overexpressed d-cofilin co-sedimented with actin filaments, suggesting that the sequestering activity of d-cofilin is weak in vivo. The overexpression of d-cofilin increased actin bundles just beneath ruffling membranes where d-cofilin was co-localized. The overexpression of d-cofilin also stimulated cell movement as well as membrane ruffling. We have demonstrated in vitro that d-cofilin transformed latticework of actin filaments cross-linked by α-actinin into bundles probably by severing the filaments. D. discoideum cofilin may sever actin filaments in vivo and induce bundling of the filaments in the presence of cross-linking proteins so as to generate contractile systems involved in membrane ruffling and cell movement.

Cell movement is a complicated phenomenon exhibited by living organisms, and is directed by interactions among proteins, membrane lipids, and extracellular matrices. Cell movement may be divided into three elementary processes; protrusion of leading edges, association and dissociation of membranes to substrata, and backward retraction of detached peripheral membranes. Actin filaments, one of cytoskeletal components, are organized into highly ordered architectures, and are involved in these processes. The architecture of actin filaments is regulated by several kinds of actin-binding proteins including cross-linking proteins, severing proteins, end-capping proteins, and monomer-sequestering proteins (Stossel et al., 1985; Pollard and Cooper, 1986; Vandekerckhove and Vancoopernolle, 1992).

Cofilin is a low molecular weight actin-modulating protein that is ubiquitously distributed among eucaryotes (Pollard, 1993; Sun et al., 1995). Cofilin-related proteins have been shown to be essential for Dictyostelium discoideum (Aizawa et al., 1995) and Saccharomyces cerevisiae (Iida et al., 1993; Moon et al., 1993) to be viable. Since both porcine cofilin and destrin, a cofilin-related protein, complement the lethality associated with yeast cofilin-null mutants, functions of cofilin-related proteins are conserved among eucaryotes (Iida et al., 1993). Cofilin-related proteins bind to monomeric actin in one-to-one molar ratio, and consequently sequesters it from self-assembly reaction in vitro (Nishida et al., 1985; Moriyama et al., 1992; Hayden et al., 1993). Cofilin-related proteins bind also to actin filaments in vitro, and occasionally severs them (Nishida et al., 1985; Cooper et al., 1986; Maciver et al., 1991A; Hawkins et al., 1993). In a variety of cells, cofilin-related proteins are localized in cellular regions which exhibit movement, such as ruffling membrane in D. discoideum cells, Acanthamoeba, and cultured fibroblastoma, and cortical actin patches in S. cerevisiae (Bamburg and Bray, 1987; Yonezawa et al., 1987; Moon et al., 1993; Quirk et al., 1993; Aizawa et al., 1995). These facts suggest the relationship of cofilin-related proteins with cell movements.

To elucidate in vivo functions of cofilin-related proteins, we examined biological phenotypes of D. discoideum cells in which the expression of D. discoideum cofilin (d-cofilin)1 was experimentally increased.

Materials and Methods

Cell Lines and Vectors

Overexpression plasmid of d-cofilin, pCOF, was constructed as follows. The d-cofilin cDNA was amplified by PCR using a synthetic oligonucleotide (AAAAATGCATCTTACAGGTATTCAGTTAGCT) and a T7 primer as

1. Abbreviation used in this paper: d-cofilin, D. discoideum cofilin.
primers from the template plasmid, pDCOF2 (Aizawa et al., 1995). The amplified cDNA was sequenced to identify a coding region. The result of sequencing was compared with that of the template. The amplified cDNA was then used for transformation experiments. The resulting plasmids were analyzed to determine the DNA content and the insertion size.

The Journal of Cell Biology, Volume 132, 1996 336

and 1-vol packed Ax2 cell pellet is 10 mg/ml. Transformed cells (2 x 10^7 cells) were grown on a 9 cm dish. After starvation for 1 h at 25°C, the plate completely. The complete lysis was checked under a microscope. Then, 10 ml of lysis buffer containing 0.5% NP-40 was added gently onto the plate, and was incubated at 22°C for 5 min to lyse all the cells on the plate. An aliquot (500 μl) of the suspension was added to 9-vol lysis buffer to lyse all the cells. The supernatant fraction (6 ml) was adjusted to 40% saturation of ammonium sulfate, and centrifuged at 10,000 rpm for 15 min, and the supernatant was collected and resuspended in 1 ml of MEM buffer as the 0-40% ammonium sulfate fraction.

Protein Compositions

Transformed cells (1 x 10^7 cells on a 9 cm dish) were harvested and grown in 80 ml of HLS medium with 10 μg/ml neomycin as liquid culture shaking at 22°C to a density at 1 x 10^7 cells/ml. Cells were harvested by centrifugation at 250 g for 5 min at 4°C, and the packed cells (0.6 g) were suspended with 6 ml of MEM buffer (20 mM Mes, 2 mM EGTA, 1 mM MgCl2, pH 6.85) containing protease inhibitors (1 mM PMSF, 50 μg/ml leupeptin, and 0.5% [vol/vol] aprotinin) and sonicated to prepare total protein fraction. After dialysis against MCM buffer (20 mM Mes, 0.2 mM CaCl2, 2 mM EGTA, pH 6.85) for 1 h at 22°C, the total protein concentration of the cell homogenate with 9-vol lysis buffer was determined by absorbance at 220 nm. The supernatant fraction was collected. The supernatant fraction was adjusted to 40% saturation of ammonium sulfate, and centrifuged at 10,000 rpm for 15 min, and the precipitate was collected and resuspended in 1 ml of MEM buffer as the 0-40% ammonium sulfate fraction. The supernatant was further sequentially fractionated into the 40-80% ammonium sulfate fraction and the 60-80% ammonium sulfate fraction. After dialysis against 50 mM Tris (pH 7.5), the 60-80% fraction was charged onto 1 ml of DEAE-cellulose column, and the flow through fraction was collected. Protein composition of each fraction was analyzed by SDS-PAGE (Laemmli, 1970) with 10-20% gradient gel. Molecular weight markers for SDS-PAGE (Bio-Rad Laboratories, Hercules, CA) contain myosin (200 KD), β-galactosidase (116 KD), phosphorylase b (97 KD), bovine serum albumin (66 KD), ovalbumin (45 KD), carbonic anhydrase (31 KD), trypsin inhibitor (21.5 KD), and lysozyme (14.4 KD). Gels were stained with Coomassie brilliant blue R-250. Western blotting was performed as previously reported (Aizawa et al., 1995) using a rabbit antisera against d-cofilin (Aizawa et al., 1995) at 2,000× dilution and a mouse anti-actin monoclonal antibody clone C4 (ICN Biomedicals Inc., Costa Mesa, CA) at 500× dilution in blocking solution (5% skim milk in PBS), respectively. The respective antibodies, alkaline phosphatase-conjugated goat anti-rabbit IgG (Jackson ImmunoResearch Laboratories, West Grove, PA) and horse radish peroxidase-conjugated goat anti-mouse IgG were used as the second antibodies, at 2,000× dilution. Reacted proteins were visualized by adding nitro blue tetrazolium chloride and 5-bromo-4-chloro-3-indolyl phosphate for staining d-cofilin, and H2O2, 4-chloronaphthol, and H2O2, 4-chloronaphthol, and H2O2, 4-chloronaphthol, and 4-dianisidine for staining actin, as substrates.

Subcellular Fractionation

Transformed cells (2 x 10^5 cells) were grown on a 9 cm dish. After starvation for 1 h at 22°C in MEM buffer (20 mM Mes, 0.2 mM CaCl2, 2 mM MgCl2, pH 6.85), the cells were washed twice on the plate by lysis buffer (20 mM Mes, 15 mM KCl, 2 mM MgCl2, 5 mM EGTA, 1 mM DTT, 0.5% NP-40, the protease inhibitors, pH 6.85) at 22°C. After a complete removal of the lysis buffer, 0.5 ml of lysis buffer containing 0.5% NP-40 was added gently onto the plate, and was incubated at 22°C for 5 min to lyse all the cells on the plate completely. The complete lysis was checked under microscopy. The lysate was collected by cell scrapers as the total fraction. The lysate was centrifuged at 400,000 g for 30 min, and the supernatant and precipitates were collected separately. The amounts of d-cofilin and actin were measured by densitometry after SDS-PAGE of each fraction. We calculated intracellular concentrations of the two proteins from the data that the total protein concentration of cell homogenate with 9-vol lysis buffer and 1-vol packed Ax2 cell pellet is 10 mg/ml.

Cell Staining

Cells grown on a glass cover slip were starved in MCM buffer for 1 h, and then fixed in MCM buffer containing 3.7% formaldehyde. For single staining of actin filaments, the fixed cells were incubated in PBS containing 1 U/ml rhodamine phalloidin (Molecular Probe, Eugene, OR) for 1 h at 25°C. For double staining of actin filaments and plasma membrane, cells were fixed and stained by rhodamine phalloidin as above, and then incubated with 800× diluted DII stock solution (3 mg/ml in ETOH) in PBS for 30 min at 25°C. For double staining of d-cofilin and actin filaments, the fixed cells were further permeabilized by incubation in ETOH containing 1% formaldehyde at −15°C for 5 min. Then the cells were incubated sequentially in the blocking solution (10% goat serum in PBS) for 20 min at 25°C, 100× diluted anti-d-cofilin antiserum for 16 h at 4°C, 100× diluted fluorescein-conjugated goat anti-rabbit IgG antibody for 2 h at 25°C, and 1 μl/ml rhodamine phalloidin in PBS for 1 h at 25°C. The stained cells were observed and recorded under a confocal laser scanning microscopy MRC600 (Bio-Rad Laboratories, Tokyo, Japan) equipped with an Argon ion (25 mW) and Helium/Neon (0.3 mW) dual laser system, Nikon optiphot-2.2, and a Nikon Plan Apo60 oil immersion objective (Nikon Co., Tokyo, Japan), and under a Zeiss Axioint 135 equipped with an Plan-Apochromat 63× oil immersion objective (Carl Zeiss, Oberkochen, Germany).

Protein Purification

Rabbit muscle actin was prepared as described (Spudich and Watt, 1971), and further purified as described before (Aizawa et al., 1995). D. discoideum cofilin was purified from Ax2 cells as described (Condeelis et al., 1982) except for the use of HiLoadTM 16/60 Superdex TM 200 purification grade gel filtration column (Pharmacia LKB Biotechnology, Uppsala, Sweden) equilibrated with 10 mM Mes, 1 mM EGTA, pH 6.9 instead of Bio-Rad A15M gel filtration open column.

Cell Movement

Transformed cells were grown on a 35 mm dish at a density of 1 x 10^5 cells/dish for overnight in HL5 medium containing 10 μg/ml neomycin. After removal of the medium, the cells were suspended in 1 ml of MCM buffer by gentle pipetting. An aliquot (50 μl) of the suspension was added to a Falcon 35 mm tissue culture dish (Becton Dickinson Co., Mountain View, CA) containing 2 ml of MCM buffer and the dish was set at 22°C for 30 min so that the cells attached to the substrate. The movement of the cells was observed under Axioint 135 equipped with Argus 50 image analyzing computer system and STC camera C2400-08. For high-magnification observation by Nomarski DIC system, we used Plan-Apochromat 63× oil emulsion object and 35 mm glass bottom microwell dishes coated with poly-L-lysine (MatTek Corp., Ashland, MA). Original images were enhanced and subtracted with the background image in real time by Argus system and recorded sequentially at every 10 s intervals on its RAM disk. To regulate temperature of the buffer at 22°C, we placed all the observation system in a room adjusted at 22°C, and the power of light for observation was set below the range 4. For low magnification observation under light microscopy, we used Plan-Neofluar 20× objective and a Falcon tissue culture dish uncocated or coated with various materials. Original microscopic images were enhanced, subtracted with the background image, sliced, and negatively polarized in real time, and recorded sequentially on a time-lapse 8-mm video recorder (Mitsubishi, Tokyo, Japan).

Making movies and parameter calculations of cell movement were automatically performed using computer program DIAS (Softech Inc., Oakdale, LA) from the recorded 8 mm video tapes.

Results

Preparation and Characterization of Cells

Overexpressing d-Cofilin

To overexpress d-cofilin in D. discoideum Ax2 cells, we used an extrachromosomal plasmid pBIG as a shuttle vector, whose copy number is ~150 in a single cell (Firtel et al., 1985). Since the pBIG contains a neomycin resistant gene, cells carrying the plasmid were selectively cloned in HL5 medium containing 10 μg/ml neomycin (Uyeda et al., 1994). The d-cofilin gene, DCOFI1 was first introduced together with actin 8 terminator into pBIG. Transfectants

The Journal of Cell Biology, Volume 132, 1996 336
carrying the plasmid did not overproduce d-cofilin, however (data not shown). Next, we introduced actin 15 promoter, d-cofilin cDNA, and actin 8 terminator into pBIG. The resultant plasmid, pCOF, successfully overexpressed d-cofilin in Ax2 cells (Fig. 1). When 10 μg plasmid was introduced into vegetatively growing Ax2 cells (1 × 10^7 cells) by electroporation, more than 100 colonies were reproducibly formed after neomycin selection. 2 wk after electroporation, all colonies were harvested and used in further experiments. The content of d-cofilin in cells carrying pCOF was seven times higher than that in cells carrying the control plasmid pBIG (Fig. 1, A and C). We found that the content of a 42-kD protein in cells carrying pCOF also increased threefold compared to that in the control cells (Fig. 1 A). The 42-kD protein was identified as actin on the basis of immunoreactivity with a monoclonal anti-actin antibody (Fig. 1 B) and binding to DNase-I column (data not shown). The contents of proteins other than d-cofilin and actin did not differ (Fig. 1 A, lanes 3–12). The expression level of d-cofilin in the above transformants continued high for 3 wk after electroporation, and gradually decreased thereafter to the control level (data not shown). The expression of actin in cells carrying pCOF decreased during passages for several weeks as that of d-cofilin decreased. Thus, we performed all the experiments in this study within 3 wk after electroporation.

Starvation of the overexpressing cells on a filter induced

---

**Figure 1.** Protein composition of control and overexpressor cells. (A) SDS-PAGE of fractions prepared from control and overexpressor cells. (B) Western blotting of fractions using anti-actin antibody. (C) Western blotting of fractions using anti-d-cofilin antiserum. Lanes 1, 3, 5, 7, 9, and 11; fractions prepared from control cells. Lanes 2, 4, 6, 8, 10, and 12; fractions prepared from overexpressor. Lanes 1 and 2, total protein; lanes 3 and 4, ultracentrifuged supernatant; lanes 5 and 6, 0–40% ammonium sulfate fraction; lanes 7 and 8, 40–60% ammonium sulfate fraction; lanes 9 and 10, 60–80% ammonium sulfate fraction; lanes 11 and 12, DEAE flow-through fraction of 60–80% ammonium sulfate fraction. M, marker proteins.

**Figure 2.** Quantitative analysis of d-cofilin and actin in control and overexpressor cells. Total protein fraction was prepared from control and overexpressor cells by lysis buffer containing 0.5% NP-40 as described in Materials and Methods. The fraction was further fractionated by ultracentrifugation into precipitates and supernatant. The amounts of d-cofilin (A) and actin (B) in each fraction were determined by SDS-PAGE followed by densitometric analysis. The concentrations of d-cofilin and actin which were produced by pCOF were also calculated (C). The mean value of three experiments were presented.
normal differentiation into aggregation stage and mature
fruiting body at 5.5 and 22 h, respectively. Starved control
cells differentiated into the above two stages at 6 and 24 h,
respectively. The content of d-cofilin and that of actin in
the overexpressing cells decreased to 30% and 10% of the
vegetative levels at 6 and 16 h, respectively, after the in-
duction of differentiation. For this reason, we performed
all the experiments using cells in MCM buffer solution
within 3 h. No significant difference was detected between
doubling times of the overexpressing cells (17 ± 0.5 h) and
control cells (18 ± 0.5 h) when they grew on a dish in HL5
medium containing 10 μg/ml neomycin. Overexpressing
cells and control cells did not differ in phagocytic activity
with microbeads (data not shown).

Figure 3. Intracellular distribution of actin filaments in
flattened control and overexpressor cells. Cells were placed
on a glass coverslip in MCM buffer for 1 h before fixation.
The fixed cells were stained with rhodamine phalloidin to
visualize actin filaments. Optically sectioned images were re-
corded by laser scanning confocal microscopy at 0.5-μm
intervals along the vertical axis. Reconstituted figures by
projecting all the Z-series of two typical flattened cells were
represented here for control (A and B) and overexpressor
(C and D), respectively. Bars: (B) 5 μm (D) 10 μm.

Figure 4. Double-staining of overexpressor for actin filaments and plasma membrane. Overexpressor cells were grown and fixed as in
Fig. 3. The cells were doubly stained by rhodamine phalloidin and Dil to visualize actin filaments and plasma membrane, respectively.
Observation was performed as in Fig. 3. Reconstituted figures from optical sections for actin filaments (A), plasma membrane (B), and
merger of A and B (C) are represented. Bar, 5 μm.
Figure 5. Intracellular distribution of actin filaments in actively moving control and overexpressor cells. Control cells (A and B) and overexpressor (C and D) were grown, fixed, and stained with rhodamine phalloidin as in Fig. 3. Observation was performed using Nomarski differential microscopy (A and C) and fluorescent microscopy (B and D). Asterisks indicate actin latticework on contact sites with the substratum. Arrows indicate actin bundles under plasma membrane. Bar, 20 μm.

Effect of d-Cofilin Overexpression on Actin Assembly in Cells

The total concentration of d-cofilin and that of actin in cells carrying pCOF were both approximately 600 μM higher than those in control cells (Fig. 2). Cell lysates were subjected to centrifugation so as to recover polymerized actin and unpolymerized actin as precipitated and supernatant fractions, respectively. Fig. 2 C showed that 83% of overexpressed d-cofilin and 90% of co-overproduced actin were recovered in precipitated fractions, indicating that the overexpression of d-cofilin co-overproduced actin filaments but not actin monomers in D. discoideum cells.

Effect of d-Cofilin Overexpression on Actin Architectures in Cells

Actin architectures of cells were visualized with rhodamine phalloidin by fluorescence microscopy. In control cells, actin filaments were observed predominantly in the regions of peripheral ruffling membranes (Fig. 3, A and B). In cells overexpressing d-cofilin and actin filaments, numerous bundles of actin filaments just under the dorsal plasma membranes were observed (Fig. 3, C and D). Double staining of actin filaments with rhodamine phalloidin and plasma membranes with dif in a single flattened cell clearly revealed that the overexpression of d-cofilin significantly stimulated membrane ruffling in dorsal cell surfaces which appeared to be associated with bundles of actin filaments (Fig. 4).

Next, we examined actin architectures in cells with elongated cell shapes which were actively moving. In control cells, we observed at least three types of phalloidin staining; first, relatively strong staining at the ventral adhesion

Figure 6. Double staining of overexpressor for actin filaments and d-cofilin. Overexpressor cells were grown and fixed as in Fig. 3. The cells were doubly stained by rhodamine phalloidin and anti-d-cofilin antibodies to visualize actin filaments (A, C, E, and G) and d-cofilin (B, D, F, and H), respectively. Observations were performed as in Fig. 3. Reconstituted figures by projecting all the Z series are shown at the top (A and B). Sectioned images at 3.0 μm (C and D), 1.5 μm (E and F), 0 μm (G and H) from the glass substrate are also shown. Bar, 20 μm.
Figure 7. Reconstitution of actin bundles from actin filaments, α-actinin, and d-cofilin in vitro. Purified rabbit muscle actin (10 μM) and D. discoideum α-actinin (0.5 μM) were mixed in the absence (A) and presence (B) of d-cofilin (1 μM) at 4°C. The final condition of reaction mixture was 10 mM Mes, 50 mM KCl, 1 mM EGTA, 0.5 mM ATP, 1 mM MgCl₂, 0.1 μM rhodamine phalloidin, pH 6.85. The mixture was incubated at 25°C for 2 h in a small cuvette made of a slide glass and a cover glass with 0.15-mm-thick spacers. The actin architectures reconstituted in the cuvette were observed under a Zeiss Axiophot fluorescent microscope, and the images were recorded in a computer image analyzer system through a SIT camera. In order to demonstrate clearly the actin bundles, we subtracted a background fluorescent image corresponding to the fine actin filaments which distributed uniformly all over the field from the original images of actin architectures, and present here the resultant figures. Bar, 200 μm.

Enhancement of Actin Filament Bundling by d-Cofilin In Vitro

It has been demonstrated that actin latticework was transformed into bundles when actin filaments cross-linked by α-actinin in the latticework were severed by treatments with actophorin, a cofilin-related protein in Acanthamoeba in vitro (Maciver et al., 1991b). We performed an analogous experiment using d-cofilin as a possible severing factor (Fig. 7). Actin latticework was reconstituted from 10 μM actin and 0.5 μM D. discoideum α-actinin (Fig. 7 A). The latticework was not brightly stained with rhodamine phalloidin because each actin filament was thin and, therefore, stained under a threshold level to be detected. When the latticework was treated with 1 μM d-cofilin, the bundles of actin filaments brightly stained with rhodamine phalloidin were induced (Fig. 7 B).

Amoeboid Movement

During the course of selection for cells containing plasmids, pCOF and pBIG, we realized that colonies of cells overexpressing d-cofilin were generally larger than those of the control cells. At day 7 and 11 after plating, the overexpressing cells formed approximately 1.6-times larger colonies in radius than the control cells (Fig. 8). Doubling time of the overexpressing cells and that of the control cells were 17 ± 0.5 and 18 ± 0.5 h, respectively. It seems that this difference, if any, in doubling time does not account for difference in colony size. Since the radius of a colony is a function of growth rate and cell motility, we speculated that the overexpressing cells move faster than the control cells. To confirm this, we decided to measure cell movement directly by real-time observation using a computer-digitated video-microscopy.

Although there is some deviation in the motility among cells, we found that cells overexpressing d-cofilin moved significantly faster than the control cells. Presented are sequential photographs of examples; a relatively fast moving cell of the control (Fig. 9 A) and one of the overexpressing cells (Fig. 9 B), taken at an interval of 20 s. The overexpressing cells developed a larger lamellipodium in front of the cell (Fig. 9, 20 s) and more rugged dorsal plasma membrane (Fig. 9, 40 s) than the control cell. For statistic treatment of motility data, we used an automatic image analyzing computer system combined with a microscope equipped with a low magnification objective. With the control and overexpressing cell types, we observed about 25 cells in each field at an interval of 3 min for 30 min (Fig. 10, A and B).
Figure 8. Colony formation of control and overexpressor cells. *D. discoideum* Ax2 cells were transformed by electroporation of plasmid vectors pBIG for control and pCOF for overexpression of d-cofilin. After electroporation, cells were grown on a dish in HL5 medium containing 10 μg/ml neomycin in order to screen cells carrying pBIG or pCOF. During the screening of transformed cells, the cells carrying the plasmids proliferated well and formed colonies, while the other Ax2 cells died. We present here photographs of typical colony of cells carrying pBIG (A and C) and pCOF (B and D) at day 7 (A and B) and at day 11 (C and D). (E) Mean radius of control and overexpressor colonies at days 7 and 11. Bar, 500 μm.
After the automatic calculation of centroid position (Fig. 10, C and D), histograms of cell frequency were made as a function of cell motility after collecting data in four independent experiments (Fig. 10, E and F). The results shown in the histograms clearly indicate that the overexpression of d-cofilin enhanced cell movement. Mean speeds of the control and overexpressing cells were calculated to be 1.09 and 1.86 μm/min, respectively. Even when the surfaces of dishes were coated with poly-L-lysine, gelatin, fibronectin, or collagen, the mean speed of the overexpressing cells (1.62, 1.16, 1.14, and 1.17 μm/min, respectively) was always about two times faster than that of the control cells (0.95, 0.51, 0.60, and 0.74 μm/min, respectively). This indicates that the enhancement of cell movement by the overexpression of d-cofilin is independent of substrate properties which affect cell motility.

**Discussion**

In this study, to address the question of what is (are) the in vivo function(s) of cofilin, we have analyzed altered properties associated with *D. discoideum* cells in which d-cofilin was overexpressed. We found that the overexpression of d-cofilin caused (a) an increase in the amount of actin filaments but not in the amount of unpolymerized actin (Fig. 2), (b) an appearance of ruffling membranes associated with cytoplasmic actin bundles (Figs. 3–7), and (c) an enhancement of amoeboid movement (Figs. 8 and 9).

How did d-cofilin overexpression cause an increase in the amount of actin filaments? It was demonstrated that a cofilin-related protein protected actin molecule from denaturation by EDTA in vitro (Hayden et al., 1993). Thus, it is possible that the overexpression of d-cofilin stabilizes actin molecules and consequently causes the accumulation of actin in the cells. Since the contribution of d-cofilin to sequestering monomeric actin is small as described below, the amount of polymerized actin is considered to increase as the amount of total actin increases. Another possibility is that severing activity of d-cofilin may cause an increase in the number of free ends of actin filaments where monomeric actin molecules are preferentially polymerized, and that lowering the concentration of free monomeric actin might enhance the synthesis of actin. The latter possibility may be supported by the recent report that the underexpression of capping protein, which increased the number of free ends of actin filaments, resulted in increases of cellular contents of both total and polymerized actin presumably because of the same mechanism as in the case of the overexpression of d-cofilin (Hug et al., 1995).

We previously reported that 9 μM d-cofilin depolymerized and sequestered only 1 μM actin in the presence of 3.2 μM total actin at pH 6.8 in vitro (Aizawa et al., 1995). Several reports have also demonstrated that about a 10 mol excess of cofilin-related proteins is needed to seques-
ter actin monomer at neutral pH (Nishida et al., 1984; Moriyama et al., 1992; Hawkins et al., 1993; Hayden et al., 1993). This weak sequestering activity of cofilin-related proteins well explains the fact that the overexpression of 600 μM d-cofilin by pCOF only slightly increased the concentration of sequestered actin from 120 to 180 μM while it greatly increased the concentration of actin filaments from 180 to 720 μM in D. discoideum cells (Fig. 2). Since the concentration of d-cofilin in Ax2 cells was estimated to be 100 μM, d-cofilin may sequester ca. 10 μM actin in the cells. Since the total concentration of sequestered actin in Ax2 cells was calculated to be ca. 120 μM, 110 μM actin should be sequestered by other proteins than d-cofilin such as profilin. We concluded that d-cofilin is not a major actin-sequestering protein in D. discoideum cells.

We showed that the overexpression of d-cofilin enhanced actin bundles in D. discoideum cells (Figs. 3–6). It was also reported that microinjection or transient overexpression of cofilin-related proteins induced actin bundles in mammalian cultured cells (Nagaoka et al., 1995; Moriyama et al., in press). These results suggest that cofilin-related proteins trigger the formation of actin bundles in cells although purified cofilin-related proteins themselves do not have any activity to bundle actin filaments in vitro. Porcine destrin, Acanthamoeba actophorin, and human actin depolymerizing factor, all of which are cofilin-related proteins, have an activity to sever actin filaments in vitro (Nishida et al., 1985; Cooper et al., 1986; Maciver et al., 1991a; Hawkins et al., 1993). It has been shown that actophorin re-organized latticework consisting of actin filaments and a cross-linking protein, into actin bundles by severing actin filaments in vitro (Maciver et al., 1991b). We also observed the lattice-bundle transition of actin filaments by d-cofilin in vitro (Fig. 6). These in vitro results suggest that the formation of actin bundles in the overexpressing cells is attributed to the sequestering activity of cofilin-related proteins.

Cells overexpressing d-cofilin exhibited enhanced cell motility (Figs. 9 and 10). This indicates that d-cofilin is an upstream positive regulator of cell motility. Very recently, phosphorylation of cofilin-related proteins at their Ser-3 residue has been shown to inactivate the protein in actin binding (Agnew et al., 1995; Moriyama et al., in press). We observed that the overexpressed d-cofilin was not significantly phosphorylated and that the overexpression of d-cofilin mutant containing Glu-3 instead of Ser-3 did not overproduce actin bundles, ruffling membranes, or cell motility in D. discoideum (data not shown). A body of evidence has been accumulated that various cellular activations accompany dephosphorylation of cofilin-related proteins and enhanced cell motilities such as stimulation of ruffling membranes (Davidson and Haslam, 1994) and secretion (Saito et al., 1994; Kanamori et al., 1995). These correlations are consistent with our conclusion that the activation of cofilin-related proteins may cause the enhancement of cell motilities. The enhancement of cell motility may be accounted for by the co-overproduction of actin filaments, since d-cofilin itself is not shown to produce mechanochemical power. The majority of actin filaments in the overexpression cells were organized into bundles under ruffling membranes (Figs. 3–6). Recently it was reported that underexpression of capping protein caused overproduction of actin filaments which were organized into actin arrays just in microspikes, but did not enhance cell motility (Hug et al., 1995). These results suggest that the enhancement of cell motility induced by the overexpression of d-cofilin is not solely attributed to the overproduction of actin filaments. We suggest that the overexpression of d-cofilin may stimulate the lattice-to-bundle transition of actin architectures, leading to cell locomotion in D. discoideum cells.

We would like to express our special thanks to Dr. K. Iida and Mr. K. Moriyama for various important suggestions and discussions on this experiment. We also thank A. Ishii, M. Yonehara, Dr. S. Matsumoto, Dr. Y. Miyata, T. Ishizawa, and K. Kimura for fruitful discussions and various technical assistance.

This work was supported in part by grants-in-aid from the Ministry of Education, Science and Culture of Japan.

Received for publication 11 September 1995 and in revised form 30 October 1995.

References

Agnew, B. J., L. S. Minamide, and J. R. Bamburg. 1995. Reactivation of phosphoactin depolymerizing factor and identification of the regulatory site. J. Biol. Chem. 270:17582–17587.

Aizawa, H., K. Sutoh, S. Tsukui, S. Kawashima, A. Ishii, and I. Yahara. 1995. Identification, characterization, and intracellular distribution of cofilin in Dictyostelium discoideum. J. Biol. Chem. 270:10923–10932.

Bamburg, J. R., and D. Bray. 1987. Distribution and cellular localization of actin depolymerizing factor. J. Cell Biol. 105:2817–2825.

Condeelis, J., S. Geoactis, and M. Vahey. 1982. Isolation of a new actin-binding protein from Dicytete6omy lactis coagulase. Cell Motil. 2:273–285.

Cooper, J. A., D. C. Cooper, R. J. Williams, and T. D. Pollard. 1986. Purification and characterization of actothen, a new 15,000 dalton actin-binding protein from Acanthamoeba castellani. J. Biol. Chem. 261:477–485.

Davidson, M. M., and R. J. Haslam. 1994. Dephosphorylation of cofilin in stimulated platelets: roles for a GTP-binding and Ca~+~. Biochem. J. 301:41–47.

Firtel, R. A., C. Silan, T. E. Ward, P. Howard, B. A. Metz, W. Nellen, and A. Jacobson. 1985. Extrachromosomal replication of shuttle vectors in Dictyostelium discoideum. Mol. Cell. Biol. 5:2541–2556.

Hawkins, M., B. Pope, S. K. Maciver, and A. G. Woods. 1993. Human actin demethylizing factor mediates a pH-sensitive destruction of actin filaments. Biochemistry. 32:9985–9993.

Hayden, S. M., P. S. Miller, A. Brauweiler, and J. R. Bamburg. 1993. Analysis of the interactions of actin demethylizing factor with F- and G-actin. Biochemistry. 32:9994–10004.

Hug, C., P. Y. Jay, I. Reddy, J. G. McNally, P. C. Bridgman, E. L. Elson, and J. A. Cooper. 1995. Capping protein levels influence actin assembly and cell motility in Dictyostelium. Cell 81:519–600.

Iida, K., K. Moriyama, S. Matsumoto, H. Kawasaki, E. Nishida, and I. Yahara. 1993. Isolation of a yeast essential gene, COF1, that encodes a homologue of mammalian cofilin, a low-Mr actin-binding and depolymerizing protein. Gene 126:115–120.

Kanamori, T., T. Hayakawa, M. Suzuki, and K. Titani. 1995. Identification of two 17-kDa rat parotid gland phospholipid phospholipases, subjects for dephosphorylation upon β-adrenergic stimulation, as destrin- and cofilin-like proteins. J. Biol. Chem. 270:8061–8067.

Laemmli, U. K. 1970. Cleavage of structural proteins during the assembly of the head of bacteriophage T4. Nature (Lond.). 227:680–685.

Maciver, S. K., H. G. Zot, and T. D. Pollard. 1991a. Characterization of actin filament severing by actophorin from Acanthamoeba castellani. J. Cell Biol. 115:1661–1620.

Maciver, S. K., D. H. Wachstock, W. H. Schwarz, and T. D. Pollard. 1991b. The actin filament severing protein promotes the formation of rigid bundles of actin filaments crosslinked with α-actinin. J. Cell Biol. 115:1621–1628.

Moon, A. L., P. A. Jannay, K. A. Louie, and D. G. Drubin. 1993. Cofilin is an essential component of the yeast cortical cytoskeleton. J. Cell Biol. 120:421–435.

Moriyama, K., K. Iida, and I. Yahara. 1996. Phosphorylation of Ser-3 of cofilin regulates its essential Function on Actin. Genes to Cells. In press.

Moriyama, K., N. Yonezawa, H. Sakai, I. Yahara, and E. Nishida. 1992. Mutational analysis of an actin-binding site of cofilin and characterization of chimeric proteins between cofilin and destrin. J. Biol. Chem. 267:7240–7244.

Nagaoka, R., K. Kusano, H. Abe, and T. Ohtani. 1995. Effects of cofilin on actin filamentous structures in cultured muscle cells. J. Cell Sci. 108:581–593.

Nishida, E., S. Maekawa, and H. Sakai. 1984. Cofilin, a protein in porcine brain that binds to actin filaments and inhibits their interactions with myosin and tropomyosin. Biochemistry. 23:5307–5313.

Nishida, E., E. Muneyuki, S. Maekawa, Y. Ohta, and H. Sakai. 1985. An actin-
Depolymerizing protein (destrin) from porcine kidney. Its action on F-actin containing or lacking tropomyosin. Biochemistry. 24:6624-6630.

Pollard, T. D. 1993. Actin and actin binding proteins. In Guidebook to the cytoskeletal and motor proteins. T. Kreis and R. Vale, editors. Oxford University Press, Oxford, UK. 3-11.

Pollard, T. D., and J. A. Cooper. 1986. Actin and actin-binding proteins. A critical evaluation of mechanisms and functions. Annu. Rev. Biochem. 55:987-1035.

Quirk, S., S. K. Maciver, C. Ampe, S. K. Doberstein, D. A. Kaiser, J. V. Damme, J. S. Vandekerckhove, and T. D. Pollard. 1993. Primary structure of and studies on Acanthamoeba actophorin. Biochemistry. 32:8525-8533.

Saito, T., F. Lamy, P. P. Roger, R. Lecocq, and J. E. Dumont. 1994. Characterization and identification as cofilin and destrin of two thyrotropin- and phorbol ester-regulated phosphoprotein in thyroid cells. Exp. Cell Res. 212:49-61.

Spudich, J. A., and S. Watt. 1971. The regulation of rabbit skeletal muscle contraction. I. Biochemical studies of the interaction of the tropomyosin-troponin complex with actin and the proteolytic fragments of myosin. J. Biol. Chem. 246:4866-4871.

Stossel, T. P., C. Chaponnier, R. M. Ezzel, J. H. Hartwig, P. A. Janmey, D. J. Kwiatkowski, S. E. Lind, D. B. Smith, F. S. Southwick, H. L. Yin, and K. S. Zaner. 1985. Non-muscle actin binding protein. Annu. Rev. Cell Biol. 1:353-402.

Sun, H. Q., K. Kwiatkowska, and H. L. Yin. 1995. Actin monomer binding proteins. Curr. Opin. Cell Biol. 7:102-110.

Sutoh, K. 1993. A transformation vector for Dictyostelium discoideum with a new selectable marker bsr. Plasmid. 30:150-154.

Uyeda, T. Q., K. M. Ruppel, and J. A. Spudich. 1994. Enzymatic activities correlate with chimeric substitution at the actin-binding face of myosin. Science (Wash. DC). 368:567-569.

Vandekerckhove, J., and K. Vancompernolle. 1992. Structural relationships of actin-binding proteins. Curr. Opin. Cell Biol. 4:36-42.

Yonezawa, N., E. Nishida, S. Koyasu, S. Maekawa, Y. Ohta, I. Yahara, and H. Sakai. 1987. Distribution among tissues and intracellular localization of cofilin, a 21kD actin-binding protein. Cell Struct. Funct. 12:443-452.