Identification and Isolation of Actin from *Neurospora crassa*

By LEN SIKORA and GEORGE A. MARZLUF*

Department of Biochemistry, Ohio State University, Columbus, Ohio 43210, U.S.A.

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Crude cell extracts of *Neurospora crassa* contained an abundant protein that was identified as actin by a number of criteria. The protein, either in cell extracts or in pure form, co-migrated with rabbit skeletal muscle actin in polyacrylamide gels. The *N. crassa* actin was purified by DEAE-cellulose and DNAase I–Sepharose chromatography and had the expected property of inhibiting DNAase I activity. Although *N. crassa* actin could polymerize and depolymerize, purification based entirely on this characteristic was ineffective. The actin was susceptible to proteolytic degradation, and under certain conditions, a breakdown product of defined size was observed.

**INTRODUCTION**

Muscle cell actin has been the focus of numerous studies due to its involvement in the contractile apparatus. Actin is also a major protein in many non-muscle tissues and cell types of eukaryotic organisms (Korn, 1978; Pollard & Weihing, 1974). In non-muscle cells actin is presumed to play a role in the functioning of the cytoskeleton in addition to participating in a number of cellular contractile processes. Non-muscle actins have many features in common and share certain biochemical properties with skeletal muscle actin. These properties include amino acid composition, molecular weight, the ability to form F-actin and to activate Mg$^{2+}$-dependent myosin ATPase activity and the ability to inhibit deoxyribonuclease I (DNAase I). Purification procedures based upon these biochemical characteristics have facilitated the isolation of actin from several lower eukaryotic organisms, most notably *Dictyostelium*, *Physarum* and *Saccharomyces* (Hatano & Oosawa, 1966; Henney & Izadpanah, 1979; Kotelliansky et al., 1979; Uyemura et al., 1978).

In *Dictyostelium discoideum*, actin is an abundant protein and it is developmentally regulated in differentiating cells (Alton & Lodish, 1977; Margolskee & Lodish, 1980; Spudich, 1974). Recombinant DNA techniques have indicated that this organism possesses multiple actin genes as well as different size classes of actin mRNA (Bender et al., 1978; Kindle & Firtel, 1978; McKeown et al., 1978).

Allen & Sussman (1978) demonstrated by electron microscopy that crude *Neurospora crassa* extracts display a characteristic 'arrowhead' configuration after incubation with heavy meromyosin, suggesting the presence of actin. Here, we present biochemical data which indicates that *N. crassa* cells possess a protein with properties characteristic of actin.

**METHODS**

*Organism and growth conditions.* Wild-type *Neurospora crassa* strain 74 OR 231 A was obtained from the Fungal Genetics Stock Center, Arcata, Calif., U.S.A. Cultures were grown for 18–20 h in Vogel's Medium N (Vogel, 1956) supplemented with 1-5% (w/v) sucrose, and then harvested and washed with distilled water. The mycelium was either extracted with acetone (Munkres, 1962) or used directly for actin isolation.

*Cell extract preparation.* An acetone powder (10 g) of *N. crassa* mycelium was extracted as described by Spudich & Watt (1971) except all solutions contained 1 mm-phenylmethylsulphonyl fluoride (PMSF) and...
DEAE-cellulose chromatography. The column (35 × 2 cm) of DEAE-cellulose (Whatman DE-52) was equilibrated with 10 mM-triethanolamine, 5 mM-sodium pyrophosphate, 20 mM-KCl, 50 μM-ATP, pH 8 (Henney & Izadpanah, 1979); all procedures were carried out at 4 °C. An equal volume of 10 mM-triethanolamine, 1 mM-2-mercaptoethanol, 1 mM-PMSF, 1 mM-benzamidine, 50 μM-MgCl₂, pH 8 was added to the cell extract prior to its application to the DEAE-cellulose column. After the sample was applied, the column was washed with 120 ml 10 mM-triethanolamine, 5 mM-sodium pyrophosphate, 20 mM-KCl 50 μM-MgCl₂, pH 8. The column was eluted with a 11 linear gradient of 0-02-1.0 M KCl in the starting buffer. The flow rate was maintained at 50 ml h⁻¹ and fractions were collected and monitored at 290 nm. Fractions that inhibited DNAase I activity (see below) were pooled and applied to a DNAase I-Sepharose affinity column.

DNAase I–Sepharose chromatography. DNAase I–Sepharose was prepared essentially according to Dhermy et al. (1978). CNBr-activated Sepharose 4B (6 g; Pharmacia) was incubated with 40 mg DNAase I dissolved in 30 ml 0-1 m-NaHCO₃, 0-5 m-NaCl and incubated at room temperature for 2 h. The DNAase I column was washed twice with 0-1 m-NaHCO₃, 0.5 m-NaCl and then treated with 1 m-ethanolamine, pH 8 for 2 h. Following washes with 0-1 m-sodium acetate, 1 m-NaCl, pH 4 and 0-1 m-sodium borate, 1 m-NaCl, pH 8, the DNAase I–Sepharose was packed into a column (1.5 × 10 cm). The column was washed with 200 ml 0-01 m-Tris, 0-001 m-CaCl₂, pH 7-5 and operated at 4 °C. Before fractions from the DEAE-cellulose column were applied to this DNAase I affinity column, they were mixed with an equal volume of buffer A (0-5 m-sodium acetate, 1 mM-CaCl₂, 15% (v/v) glycerol, pH 6-5). After application of the sample, the DNAase I column was washed with buffer A (60 ml) until the A₆₅₀ of the eluate was reduced to the background reading. The column was then eluted with 32 ml buffer A containing 0-75 m-guanidinium chloride (GuHCl), followed by 40 ml buffer A containing 3 m-GuHCl. The fractions eluted with GuHCl were pooled and dialysed for 24 h at 4 °C against distilled water containing 1 mM-PMSF and 1 mM-benzamidine. Following dialysis, the sample was lyophilized.

DNAase I inhibition assay. Two methods were used to assay for DNAase I inhibition by the N. crassa actin-like protein. The first method was essentially as described by Kotelliansky et al. (1979) except that 1 μg DNAase I was used and the total assay volume was 10 ml. For the second method, DNA-methyl green was used as the substrate for DNAase I. DNA-methyl green was dissolved in 100 mM-Tris, 4 mM-MgCl₂, 1.7 mM-CaCl₂, pH 7-5 to a final concentration of 0-25 mg ml⁻¹ and was stored at 4 °C prior to use. DNAase I was dissolved in 0-15 m-NaCl to a final concentration of 100 μg ml⁻¹ and stored frozen. DNAase I solution (10 μl) and up to 0-5 ml of an actin-containing sample were added to cuvettes and incubated at room temperature for 0-5 h. Then 0-3 ml of the DNA-methyl green solution was added and incubation was continued for 8 h. After incubation, each cuvette was brought to 1 ml with 100 mM-Tris, 4 mM-MgCl₂, 1.7 mM-CaCl₂, pH 7-5. A blank containing only DNA-methyl green and a control containing both DNA-methyl green and DNAase I were incubated in parallel with the samples. Inhibition of DNAase I by actin was calculated as follows:

\[
\text{Percentage inhibition of DNAase I} = 100 \times \frac{(A_{650} \text{ blank} - A_{650} \text{ sample})}{(A_{650} \text{ blank} - A_{650} \text{ control})}
\]

Sodium dodecyl sulphate (SDS)–polyacrylamide gel electrophoresis. This was performed according to O'Farrell (1975) with 12.5% gels using a stacking gel in a vertical slab gel apparatus (15 × 15 cm). Molecular weights assigned to the standard proteins were: bovine serum albumin (68,000); actin (42,000); DNAase I (31,000); myoglobin (18,000).

Chemicals. Rabbit skeletal muscle actin was from Worthington. All other reagents were purchased from Sigma.

RESULTS

In attempts to determine whether extracts of *Neurospora crassa* contain an actin-like protein, cultures were processed in one of two ways. Cells were either precipitated with acetone prior to extraction of cellular proteins or used directly in the homogenization procedures. No differences in the electrophoretic patterns of proteins were observed between acetone-extracted cells and cells used directly in the homogenization procedures. For this reason, cells for actin isolation were used immediately after harvesting and were extracted as described in Methods. Cells processed in buffers lacking PMSF and benzamidine showed no discernible protein patterns following electrophoresis. Thus, in all subsequent isolation steps PMSF and benzamidine were added to minimize protein degradation. The *N. crassa* cell extracts contained a protein which co-migrated during electrophoresis with rabbit skeletal muscle...
Neurospora crassa actin

Fig. 1. SDS-polyacrylamide gel electrophoresis of fractions obtained during actin purification. Lanes 1 and 5, protein standards – bovine serum albumin, actin, DNAase and myoglobin (20 μg each protein). Lane 2, N. crassa protein sample from the DNAase I-Sepharose column (25 μg protein). Lane 3, fraction 54 from the DEAE-cellulose column (25 μg protein). Lane 4, total soluble proteins from N. crassa mycelium (125 μg protein). The arrow identifies actin.

Fig. 2. DEAE-cellulose chromatography of N. crassa crude cell extract. The column was eluted as described in the Methods. DNAase I inhibition was determined using the DNA-methyl green assay (□). Elution of protein was monitored by absorbance at 290 nm (●). Conductivity was used to determine the KCl concentration (△).

actin (Fig. 1). Furthermore, this protein represented approximately 2% of the total soluble proteins in cell extracts as judged by densitometric scans of stained polyacrylamide gels.

Efforts to purify actin by repeated rounds of polymerization and depolymerization met with limited success. After one round of polymerization and depolymerization, a protein co-migrating with authentic actin was enriched compared with the amount found in the
original whole cell extract (data not shown). However, the total spectrum of proteins found in the cell extract was also present in the depolymerized sample, and the low molecular weight proteins appeared to be present in greater amounts than the higher molecular weight proteins. Following a second round of polymerization, the proportion of proteins co-migrating in gels with authentic actin was very much decreased, but a new band greater in molecular weight than actin was observed. This band most likely represented a dimeric actin form. The extended dialysis necessary to depolymerize F-actin may have resulted in protein degradation and/or incomplete depolymerization, accounting for the loss of the actin-like protein. Similar results were obtained even when PMSF and benzamidine were included in the polymerization and depolymerization buffers.

Ion-exchange chromatography using DEAE-cellulose has proven useful in isolating actin from other organisms (Gordon et al., 1976; Henney & Izadpanah, 1979; Kotelliansky et al., 1979). When an *N. crassa* cell extract was chromatographed on DEAE-cellulose and eluted with a KCl gradient, the profile shown in Fig. 2 was obtained. DNAase I-inhibiting activity was found in the latter half of the first large protein peak and consistently eluted between 0-10 and 0.35 M-KCl. When the crude extract was chromatographed on DEAE-cellulose with an entirely different buffer system and eluted with a shallower KCl gradient (Gordon et al., 1976) the DNAase I-inhibiting activity still eluted within the same KCl concentration range (data not shown). Furthermore, actin isolated from yeast as a control showed the same elution characteristics (data not shown). The 0-10–0.35 M-KCl eluate inhibited DNAase I in both assays used (see Methods).

A sample of each fraction from the DEAE-cellulose column was assayed for DNAase I inhibition and its protein content was examined by electrophoresis in an SDS–polyacrylamide gel. Fraction 54 (see Fig. 2) inhibited actin to the greatest extent and gel electrophoresis revealed that it contained several proteins, the major one of which co-migrated with skeletal muscle actin (Fig. 1). Densitometric scans of SDS–polyacrylamide gels of this fraction indicated that it contained 30% actin. Other fractions that inhibited DNAase I also contained this protein. The DNAase I-inhibiting fractions (47–63) from the DEAE-cellulose column were pooled and chromatographed on a DNAase I–Sepharose affinity column. A protein profile of the fractions eluted from the affinity column is shown in Fig. 3. The first GuHCl wash (0.75 M) was ineffective in eluting all of the bound protein from the column and a large amount of protein was eluted in the subsequent 3 M-GuHCl wash. The DNAase I–Sepharose column had been previously tested with rabbit skeletal muscle actin, which also eluted primarily in the 3 M-GuHCl wash (data not shown). By integration of the areas beneath each
Table 1. DNAase I inhibition by fractions from DEAE-cellulose and DNAase I–Sepharose columns

DNAase I inhibition was estimated by the DNA-methyl green assay.

<table>
<thead>
<tr>
<th>Column fraction</th>
<th>Sample vol. (ml)</th>
<th>Protein (µg)</th>
<th>Inhibition of DNAase I* (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>DEAE-cellulose, fraction 54</td>
<td>0:10</td>
<td>1</td>
<td>100</td>
</tr>
<tr>
<td>0:50</td>
<td>5</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>DNAase I–Sepharose, 0:75 M-GuHCI eluate</td>
<td>0:01</td>
<td>0:1</td>
<td>1:2</td>
</tr>
<tr>
<td></td>
<td>0:10</td>
<td>1</td>
<td>7:2</td>
</tr>
<tr>
<td></td>
<td>0:50</td>
<td>5</td>
<td>19:1</td>
</tr>
<tr>
<td>DNAase I–Sepharose, 3 M-GuHCI eluate</td>
<td>0:01</td>
<td>0:1</td>
<td>3:6</td>
</tr>
<tr>
<td></td>
<td>0:10</td>
<td>1</td>
<td>11:3</td>
</tr>
<tr>
<td></td>
<td>0:50</td>
<td>5</td>
<td>25:6</td>
</tr>
</tbody>
</table>

* Complete inhibition of DNAase I activity occurred when 0.5 µg of rabbit skeletal muscle actin was used in the assay procedure.

of the protein peaks eluted from the DNAase I affinity column, it was estimated that 5% of the _N. crassa_ protein applied had bound to the affinity column and was eluted with GuHCl (0.75 M and 3 M). The material eluted from the DNAase I–Sepharose column by GuHCl was concentrated and examined by SDS–polyacrylamide gel electrophoresis. We could detect only a single major protein band, which co-migrated with authentic actin (Fig. 1). This protein represented at least 80% of the protein applied to the SDS–polyacrylamide gel.

Table 1 shows DNAase I inhibition results for samples eluted from the DEAE-cellulose and the DNAase I affinity columns. A sample of fraction 54 from the DEAE-cellulose column inhibited DNAase I completely. Comparable amounts of protein eluted from the affinity column by GuHCl inhibited DNAase I only poorly. Prior experiments utilizing buffers without protease inhibitors suggest that protease degradation may occur during the DNAase I affinity chromatography and the subsequent dialysis. In fact, if the dialysis time was greater than 24 h and PMSF and benzamidine concentrations were decreased, the major protein species detected by electrophoresis had a molecular weight of only 28 000, indicative of a protease attack upon actin.

In another experiment, _N. crassa_ proteins that had undergone one round of polymerization and depolymerization were applied to the DNAase I affinity column. The profile of protein eluting from this column looked identical to that in Fig. 3 except that the _A_280 values were slightly lower. Electrophoresis of the GuHCl-eluted material revealed a faint band co-migrating with authentic actin as well as three to five other bands (data not shown).

**DISCUSSION**

The presence of an actin-like protein in extracts of _N. crassa_ was previously suggested by Allen & Sussman (1978): the appearance of negatively stained filaments in extracts of _N. crassa_ after incubation with ATP, CaCl₂ and KCl was similar to actin filaments observed in other systems with the electron microscope. We have now identified a protein that has several properties in common with skeletal muscle actins and non-muscle actins. This protein both in crude cell extracts and in its purified form co-migrates with rabbit skeletal muscle actin. It elutes from DEAE-cellulose under the same ionic conditions as does yeast actin, it undergoes polymerization–depolymerization reactions and it inhibits DNAase I activity, comparable to that achieved with rabbit skeletal muscle actin. The specificity of the DNAase I interaction with actin has itself been the basis of actin purification schemes (Dhermy _et al._, 1978; Lazarides & Lindberg, 1974) and has become a major criterion in establishing a protein as
actin. We have examined in parallel *N. crassa* and yeast actin using the DEAE-cellulose chromatographic method of Gordon *et al.* (1976). In each instance, the protein profile, the DNAase I-inhibiting activity and the electrophoretic pattern of proteins were identical. When the appropriate *N. crassa* fractions from the DEAE-cellulose column were further chromatographed on DNAase I–Sepharose, a single protein was eluted by GuHCl and it co-migrated with rabbit skeletal muscle actin.

Polymerization and depolymerization was not effective in purifying *N. crassa* actin. One round of polymerization and depolymerization produced a preparation enriched for actin but seriously contaminated with other proteins (not shown). A second round of polymerization resulted in losses of large amounts of protein, possibly due to an ineffective polymerization or inadequate depolymerization. Similar results have been observed in other systems (Henney & Izadpanah, 1979; Reichstein & Korn, 1979).

The identification of actin as an abundant protein in *N. crassa* is not surprising in view of its presumed functions in eukaryotic organisms. Assuming that rabbit skeletal muscle actin and *N. crassa* actin inhibit DNAase I activity equally, we estimate that actin comprises 1–2% of *N. crassa* soluble proteins. This figure agrees well with results of densitometric scans of the SDS–polyacrylamide gels containing total *N. crassa* soluble proteins. In *Dictyostelium discoideum*, actin normally comprises 8% of cellular proteins and can increase to 25% in differentiating cells (Alton & Lodish, 1977). In the experiments reported here, we isolated actin from *N. crassa* cells grown for 18–20 h, when they were in exponential, vegetative growth. It is possible that in other developmental stages, such as early conidial formation or during spore germination, actin may occur in greater amounts.

Actin eluted from the DNAase I affinity column inhibited DNAase I poorly. Poor DNAase I inhibition by actin after its elution from a DNAase I column has been reported by Henney & Izadpanah (1979). Denaturation of the actin most likely occurred due to the GuHCl used for its elution from the affinity column. We did not recover large amounts of actin from the DNAase I affinity column in the absence of protease inhibitors. Attempts to isolate actin from *N. crassa* mycelium were unsuccessful unless PMSF and benzamidine were added to all buffers employed in the isolation. Moreover, when the protease inhibitors were not added to the eluant of the DNAase I affinity column no actin was detected, but we found instead a single protein band of increased mobility (mol. wt 28 000), indicating proteolysis. The presence of an altered actin band of defined molecular weight may mean that protease attack is at least initially confined to a specific region of the actin molecule. The generation of actin-like artifacts by possible proteolytic attack on the actin molecule has been discussed by Water *et al.* (1980). Protease problems with *N. crassa* extracts have been recognized by other workers (Gaetner & Cole, 1976, 1977). The 'snowflake' mutant of *N. crassa* appears to contain abnormally large amounts of actin-like protein (Allen & Sussman, 1978). Future studies with this mutant may provide valuable insights into the regulation of the actin gene and the synthesis of the actin protein.

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**REFERENCES**


