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Role of Esterase gp70 and Its Influence on Growth and Development of *Dictyostelium discoideum*

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Role of Esterase gp70 and Its Influence on Growth and Development of Dictyostelium discoideum

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Abstract
Gp70 is an esterase originally called crystal protein because of its presence in crystalline structures in aggregation-competent Dictyostelium discoideum cells. Although postulated to break down spore coats, the function of gp70 in vivo was incompletely investigated. Our immunolocalization and biochemical studies of vegetative D. discoideum amoebae show that gp70 was recruited to phagosomes and found in lysosomes. Purified gp70 was effective at hydrolyzing naphthyl substrates with acyl chains typical of lipids and lipopolysaccharides, indicating that the gp70 was involved in digesting endocytosed molecules. The activity of purified gp70 was inhibited by reductants that retarded its electrophoretic mobility and verified the presence of intramolecular disulfide bonds predicted by its amino acid sequence. Compared to wild-type cells, cells overexpressing gp70 were more phagocytically active, had shorter generation times, and produced more fruiting bodies per unit area, while cells lacking gp70 were phagocytically less active with longer doubling times, developed more slowly, and had significantly fewer fruiting bodies per unit area. Consistent with the phenotype of a disrupted metabolism, one-third of the gp70-minus cells were large and multinucleated. Together, these results indicated that despite its crystalline appearance, gp70 was an active esterase involved in both the growth and the development of D. discoideum.

Keywords: esterase, endocytosis, lysosome, phagocytosis, lipopolysaccharide

Abbreviations: BSA, bovine serum albumin; Con A, concanavalin A; DAPI, 4,6-diamidino-2-phenylindole; DTT, dithiothreitol; FITC, fluorescein isothiocyanate; GT, generation time; IgG, immunoglobulin G; LPS, lipopolysaccharide; KA, Klebsiella aerogenes; mAb, monoclonal antibody; Mes, 2-(W-morpholino)ethanesulfonic acid; Mops, 3-(2-3-morpholino)propanesulfonic acid (sodium salt); PVDF, polyvinylidene difluoride; Sorensen’s buffer, 14.6 mM KHzPO₄, 2 mM Na₂HPO₄, pH 6.1; SDS-PAGE, sodium dodecyl sulfate-polyacrylamide gel electrophoresis; TRITC, tetramethylrhodamine isothiocyanate

INTRODUCTION

By electron microscopy, crystalline bodies are observed in amoebae of the cellular slime mold Dictyostelium discoideum [1, 2]. The crystal structures become more numerous between the growth and aggregation stages, are present in mature spores, and disappear only after spore germination [2–5]. The crystals contain mainly two proteins, D2 and a 69-kDa species, which due to its abundance, was named crystal protein [6]. Because the deduced amino acid sequences of crystal protein and D2 have a high degree of sequence similarity to esterases, the vesicles were called esterosomes. Since the crystals disappear during spore germination, crystal protein was postulated to be required for spore wall degradation [6]. The exact function of crystal protein, however, has not been established.

Recently, we identified in axenically grown D. discoideum cells a 70-kDa glycoprotein, gp70, associated with detergent-insoluble cytoskeletons [7]. Amino acid analyses and antibody cross-reactivity established that gp70 is the same as crystal protein, and comparative biochemical studies of strains overexpressing or lacking gp70 showed that it has esterase activity. The apparent high levels of gp70 in log-phase, axenically grown cells indicated a function other than spore coat degradation. The goal of this work was to establish the role of gp70 in D. discoideum growth and development. Based on in vivo and in vitro analyses of gp70 overexpressor and gp70-minus strains, we propose that the esterase gp70 is involved in the digestion of endocytosed nutrients and its presence is important for normal growth and development of D. discoideum.

Portions of this work were presented at the “1999 Dictyostelium International Meeting,” Bar Harbor, Maine [8].

MATERIALS AND METHODS

Growth and development of D. discoideum. D. discoideum cells (AX2) were shaken at 190 rpm in HL5 [9] at 20°C and typically harvested at 5 × 10⁶ cells/ml. Two transformed cell lines derived from this strain, AT-K₁ which lacks gp70 (crystal protein) and AT-K₂ which overexpresses gp70 [7, 10], were grown in HL5 containing 10 μg/ml G418 (ICN Biomedicals, Inc., Aurora, OH). For axenic growth, log-phase cells were diluted to 5 × 10⁶/ml in fresh HL5 and counted every 24 h. Culture densities were monitored using a hemocytometer. Growth rates on bacteria were measured following published procedures [11]. Colony growth on bacterial lawns was accomplished by inoculating D. discoideum cells with a toothpick onto a lawn of Klebsiella aerogenes (KA)² cultivated on nutrient agar [12] and then monitoring the plaque
size over time. Fruiting bodies were counted after 5 to 7 days. NC4 (provided by Dr. M. Clarke, Oklahoma Medical Research Foundation) was grown in suspensions of KA [13]. Newly germinated cells in HL5 were obtained from spores of fruiting bodies produced from axenically grown cells starved for over 24 h.

For development studies, axenically grown log-phase cells were washed once with Sorensen’s buffer (14.6 mM KH₂PO₄, 2 mM NaH₂PO₄, pH 6.1) and resuspended to 10⁶ cells/ml. Then, 10⁶ cells were pipetted onto the center of 2% nonnutrient agar plates (100 × 20 mm) and development was monitored over time.

**Microscopy.** For electron microscopy, log-phase AX2 cells in HL5 were allowed to settle for 15 min in a 35 × 10-mm Contour dish (Lux Scientific Corp., Newbury Park, CA) before fixation at room temperature for 1 h in 3% glutaraldehyde in 100 mM phosphate buffer (pH 6.8). After rinsing with the same buffer, cells were postfixed with phosphate-buffered 1% osmium tetroxide for 1 h. After dehydration through an ethanol series (25%, 50, 75, 95%, and 100%), samples were embedded in Epon 812. Thin sections were stained with uranyl acetate and lead citrate and observed with a Philips 201 (FEI Co., Hillsboro, OR) transmission electron microscope operated at 60 keV.

For immunofluorescence microscopy, spores, newly germinated cells, or log-phase cells were washed three times with Sorensen’s buffer. After resuspending to 5 × 10⁵ cells/ml, 100 μl of cells was allowed to settle onto glass coverslips (30 min for cells or 4 h for spores). In some experiments, 5 × 10⁶ beads (4.329 μm diameter; Polysciences, Inc., Warrington, PA) were added and removed after 10 min by two washes of Sorensen’s buffer. Cells were fixed with 3.7% formaldehyde in 17 mM sodium phosphate buffer, pH 6.8, for 15 min and permeabilized for 5 min with chilled methanol (–20°C) containing 1% formaldehyde [14]. Non-specific binding was blocked with 1% bovine serum albumin (BSA) in phosphate-buffered saline (PBS; 150 mM NaCl, 10 mM sodium phosphate, pH 7.4) for a minimum of 15 min. Cells were incubated with monoclonal antibody (mAb) 80 (130-80-2) or mAb 202 (129-202-6) hybridoma supernatants [6] for 1 h at room temperature, washed three times with PBS containing 0.05% Tween 20, and then stained for 1 h with fluorescein isothiocyanate (FITC)-conjugated goat anti-mouse IgG (Pierce Chemical Co., Rockford, IL) adsorbed against glutaraldehyde-fixed AX2 [13]. Samples incubated with secondary antibody when primary antibody was omitted had no significant fluorescent signals. To identify nuclei, cells were fixed with methanol and stained with 0.5 μg/ml 4,6-diamino-2-phenylindole (DAPI; Sigma Chemical Co., St. Louis, MO) [15]. Nuclei from a total of 996 AX2 and 1534 AT-Kneg cells were counted. Coverslips were mounted in buffered Gelvatol containing 50 mg/ml 1,4-diazobicyclo-(2,2,2)octane (Aldrich Chemical Co., Milwaukee, WI) [14, 16]. Cells were viewed with a Nikon Labophot microscope (Melville, NY) equipped with bright-field 40x and 60x objectives and a 60x plan Apochromat oil objective (NA 1.4) for epifluorescence. Images were acquired with a three-color chip CCD camera (DC-330, DAGE-MTI, Inc., Michigan City, IN) and a Scion CG-7 RGB color frame grabber (Scion Corp., Frederick, MD) using Scion Image 1.62 and imported into Photoshop 5.0.

**Preparation of cell fractions and purification of gp70.** Cell lysates were prepared by sonication of 10⁷ cells/ml in 2% sodium dodecyl sulfate (SDS), 0.5 mM dithiothreitol (DTT), 2 mM ethylene glycol bis(2-aminoethyl ether)-N,N,N’,N’-tetraacetic acid (EGTA), 20 mM sodium fluoride, 0.5 mM phenylmethylsulfonyl fluoride, 1 mM benzamidine hydrochloride, and 10 mM Tris, pH 7.4. To test the esterase activity of cell lysates, 2 × 10⁷ cells (AX2 or AT-Kneg) from axenic cultures were washed and solubilized overnight at 4°C in 2 ml buffer containing 20 mM 2-(N-morpholino)ethanesulfonic acid (sodium salt; Mes), pH 6.0, 500 mM NaCl, 1 mM CaCl₂, 1 mM MgCl₂, and 2.5% Triton X-100 [7]. Lysosomes and phagosomes were prepared as described [17, 18]. Cytoskeletons were purified by extracting midlog AT-K2 cells with 1% Triton, 10 mM KCl, 10 mM EGTA, 2 mM sodium azide, 10 mM imidazole, pH 7.0 [19], in the presence of 5 mM benzamidine hydrochloride, 1 mM phenylmethylsulfonyl fluoride, and 1 mg/ml leupeptin. Cytoskeletons were solubilized overnight at 4°C in 20 mM Mes, pH 6.0, 500 mM NaCl, 1 mM CaCl₂, 1 mM MgCl₂, and 2.5% Triton X-100 [7]. The solubilized sample was centrifuged at 13,000g for 1 h at 4°C to pellet actin filaments and actin-associated proteins, leaving solubilized gp70 and two minor species (see Figure 5). The solubilized gp70 was further purified either by concanavalin A (Con A) chromatography [7] or immunofinity chromatography using mAb 80 covalently coupled to protein G beads using dimethylpyrimid-lidate (Sigma [20, 21]). Purified gp70 was quantified by densitometry of Coomassie blue-stained gels using BSA as a standard.

Protein concentrations were determined either with a filter paper dye-binding assay [22] or with the bicinchoninic acid reagent (Pierce). SDS gel electrophoresis [23] and immunoblotting [24] followed standard procedures. Hybridoma cells producing mAb 80 and mAb 202 specific for gp70 [6] were grown as previously described [7]. Antibodies were purified from hybridoma supernatants using ammonium sulfate precipitation [21] and protein G chromatography [25]. Hybridoma N2 culture supernatant to the 100-kDa subunit of the *D. discoideum* vacular H⁻/ATPase was obtained from Dr. A. Fok (University of Hawaii [26]). Gels and immunoblots were scanned with an Epson ES-1200C scanner and ScionFas-tic software. Relative amounts of protein were quantified by reflectance densitometry.

**Phagocytosis and pinocytosis assays.** Phagocytosis of tetramethylrhodamine isothiocyanate (TRITC)-labeled yeast was measured according to published procedures [27]. Trypan blue (Merck, Darmstadt, Germany; 2 mg/ml in 150 mM NaCl, 20 mM sodium citrate, pH 4.4 [28]) was used to quench the fluorescence of noningested yeast [27, 29]. Sample fluorescence was measured (Perkin-Elmer LS-5B luminescence spectrometer; Norwalk, CT) at excitation and emission wavelengths of 544 and 574 nm, respectively. Fluid-phase endocytosis (pinocytosis) was measured using FITC-dextran (Sigma) according to the method of Vogel [30] as described [7] at excitation and emission wavelengths of 470 and 520 nm, respectively.

After axenic growth, and for an equivalent number of cells, the average protein content and cell volume of AT-Kneg cells were 1.7- and 1.88-fold higher, respectively, than those of AX2 cells. The volume of packed cells was measured by centrifugation of cells into a graded tube. When transferred to bacterial suspensions, the larger axenically grown AT-Kneg cells apparently split into smaller cells, yielding a corresponding increase in cell number. Thus, after growth in bacterial suspension, the average protein content and cell volume of AT-Kneg cells were 1.17- and 1.12-fold, respectively, of AX2 cells. These differences were factored into the phagocytosis and pinocytosis data, to normalize AT-Kneg to AX2 on the basis of equal volume [27].

**Esterase assay.** Con A-purified gp70 were tested for esterase activity using a modified method of Gu and Zera [31]. Sample (4 μg; 0.2 μg/ml) or buffer (20 μl) was added to 1980 μl of 50 mM 3-(N-morpholinopropanesulfonic acid, pH 7.5 (Mops buffer), containing 0.25 mM α-naphthyl acetate (initially dissolved in ethanol; Sigma). After incubation at 30°C for 30 min, reactions were stopped with 300 μl of 0.3% Fast Blue RR Salt (Sigma) and 3.5% SDS, and their absorbance was read at 605 nm (Cary 50; Varian Instruments, Sugarland, TX). Esterase activity was expressed as micromoles of α-naphthol produced per minute per micromgram of protein. To examine the pH dependence of the esterase activity of gp70, 50 mM sodium citrate (pH 3), 50 mM sodium acetate (pH 4 or 5), 0.1 M Mes (pH 6), 0.1 M Tris (pH 9), and 25 mM carbonate-bicarbonate (pH 10) buffers [32] were used instead of Mops in the assay (Table 2). Standard curves using the same buffers were prepared in parallel. To test the effect of DTT, 15 μl of gp70 (0.2 μg/ml) was incubated with 5 μl of DTT (0.2 to 1 M) and incubated for 30 min at 4°C before addition
of the sample to the Mops buffer. The activity of purified gp70 also was tested in Mops buffer using naphthyl substrates with different acyl chains including α-naphthyl valerate (5-carbon), nonanoate (9-carbon), laurate (12-carbon), palmitate (16-carbon), and β-naphthyl myristate (14-carbon); solubilized gp70 also was active with substrates β-naphthyl acetate and p-nitrophenyl acetate (data not shown). In some experiments, Triton X-100 (final concentration 1%) was added to the Mops buffer to increase the solubility of the naphthyl substrates with long acyl chains. However, 1% Triton X-100 also inhibited by 57% the esterase activity of gp70 with α-naphthyl acetate as the substrate (average of three experiments). This percentage of inhibition was factored into the calculations of esterase activity when using the naphthyl substrates with acyl chains of 12 or more carbons (Table 1).

No significant amidase (using the three substrates l-leucme p-nitroanilide, l-phenylalanine p-nitroanilide, and glycine p-nitroamylide [33]), acid, or alkaline phosphatase activity (using the substrate p-nitrophenyl phosphate) was detected. Statistical analyses were performed with the program InStat (version 2.02A; GraphPad Software, Inc., San Diego, CA).

RESULTS

Localization of gp70

Crystalline bodies in the cytoplasm of growth-phase AX2 cells were observed by electron microscopy (Figure 1). To our knowledge, this was the first image of a crystal body shown in vegetative cells. These structures had an interlattice space of 12 nm, edge dimensions that ranged from 0.1 to 0.4 μm, and corresponded to the crystals or esterosomes observed in developing cells [1–3, 5, 6] and spores [4]. Esterosomes were shown to contain mainly two proteins, crystal protein and D2. Consistent with several previously published studies of developing cells [1–3] and spores [4], the crystals we observed in vegetative cells were without obvious membranes (Figure 1), while other studies of aggregating cells showed crystalline bodies with membranes [5, 6]. This difference may have been a consequence of the different protocols used for preparing the samples for microscopy. An alternative explanation is that the crystals were released from esterosomes into cytoplasm during different developmental stages. We recently established that a 70-kDa glycoprotein, gp70, has esterase activity and is the same as crystal protein.

In studies of phagosome maturation in *D. discoideum*, we made the observation that gp70 is recruited to the vesicles as they were processed [18]. This finding implied an interaction between esterosomes and phagosomes that was verified first by indirect immunofluorescence microscopy of cells fed latex beads (Figures 2A and 2B). Esterosomes, containing concentrated gp70, collected around phagocytosed beads. Similar aggregates of esterosomes surrounding ingested beads were seen also after longer chase times (up to 300 min) and around internalized yeast (data not shown). Additional support for an active role of gp70 in the processing of phagosome contents was its localization in NC4 cells in vesicles containing bacteria (Figures 2F and 2H). Food vacuoles that presumably arose from the fusion of small phagosomes were recognized by mAb 202, indicating that they contained gp70. Concurrently recruited with gp70 to maturing phagosomes was the vacuolar H⁺-ATPase (Figure 3A), whereas actin was removed, as ob-
served previously [18] and plotted for reference (Figure 3B). The acidification of the phagosomes by the H\(^+\)-ATPase [34] likely favored the activity of enzymes (possibly from lysosomes) including gp70, an esterase, that digest food particles.

An examination of enriched lysosome preparations demonstrated the presence of gp70 in lysosomes (Figure 4). Our Coomassie blue-stained SDS gel protein profile of enriched lysosomes was similar to that of endosomes, which are closely related to these structures [35]. Although there was no obvious difference in the intensity of the Coomassie blue-stained species at 67 kDa (the position of gp70 on 10% polyacrylamide-SDS gels) between Figure 4, lanes 1 and 2, densitometric analyses
of immunoblots revealed a 1.8- to 2.5-fold enrichment of gp70 in lysosomal preparations over lysates from axenic vegetative AX2 (Figure 4; n = 3). Further analysis of the 67-kDa species on two-dimensional gels showed that other proteins contributed to the Coomassie blue-stained species at this position on SDS gels of cell homogenates (data not shown). The lysosome preparations containing gp70 had an esterase activity of 0.015 μmol/min/μg protein (mean ± SD; n = 4) at pH 7.5 using α-naphthyl acetate.

In Vitro Esterase Activity of gp70

The esterase activity of both solubilized and Con A-purified gp70 was tested with α- or β-naphthyl substrates with different acyl chain lengths (Table 1). Gp70 was active with all the substrates, having the highest activity with α-naphthyl nonanoate. However, the activity of gp70 became less efficient with increasing length of the acyl chain, with a sixfold difference in activity between the α-naphthyl acetate and α-naphthyl stearate substrates.

The pH dependence of the esterase activity of gp70 was tested with the substrate α-naphthyl acetate. Solubilized gp70 had an activity of 0.72 ± 0.03 μmol/min/μg protein (mean ± SD; n = 9) at pH 7.5. Its activity increased 16.7% to 0.84 ± 0.05 μmol/min/μg protein (mean ± SD; n = 3) at pH 6. The esterase activity of gp70 decreased at very acidic (pH 3) or basic (pH 10) pH, with optimal activity around pH 6 (Table 2). This weak pH dependence of enzyme activity would allow gp70 to remain active in the various endocytic compartments of D. discoideum [36–39].

DTT retarded the mobility of gp70 on SDS gels (Figure 5), which supports the hypothesis that native gp70 has one to two internal disulfide bonds [6]. In the presence of DTT, Con A- or immunoaffinity-purified gp70 (Figures 5A and 5B) migrated to 67 kDa, slightly faster than the BSA of the molecular standards. In the absence of DTT, gp70 migrated to 59 kDa (on a 10% polyacrylamide gel), equal to its calculated molecular mass based on the deduced amino acid sequence [6]. Species a and b (Figure 5A, lane 3) were bound by mAb 202, indicating that they were either breakdown products of gp70 or incompletely reduced molecules since they had faster mobilities like that of gp70 in the absence of DTT (Figure 5A, lane 4). Intra-molecular disulfide bonds stabilize a more compact conformation of a protein, which usually results in a faster migration in a polyacrylamide gel than its reduced form [40]. Both the disulfide bonds and the glycosylation of gp70 [7] likely contributed to the difference between the expected 59-kDa mass calculated from the amino acid sequence and the larger relative mass of 67 kDa measured from the SDS gels. A reductant-dependent shift in the mobility of gp70 was undetected in an earlier study [7].

Table 1

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Activitya</th>
</tr>
</thead>
<tbody>
<tr>
<td>α-Naphthyl acetate (2)</td>
<td>0.72 ± 0.03</td>
</tr>
<tr>
<td>α-Naphthyl valerate (5)</td>
<td>0.73 ± 0.03</td>
</tr>
<tr>
<td>α-Naphthyl nonanoate (9)</td>
<td>1.72 ± 0.02</td>
</tr>
<tr>
<td>α-Naphthyl laurate (12)</td>
<td>0.51 ± 0.01</td>
</tr>
<tr>
<td>α-Naphthyl myristate (14)</td>
<td>0.41 ± 0.03</td>
</tr>
<tr>
<td>α-Naphthyl palmitate (16)</td>
<td>0.21 ± 0.02</td>
</tr>
<tr>
<td>α-Naphthyl stearate (18)</td>
<td>0.12 ± 0.01</td>
</tr>
</tbody>
</table>

a μmol/min/μg protein; mean ± SD, n = 3-9.

b Number of carbons.
Table 2  

<table>
<thead>
<tr>
<th>pH</th>
<th>Activity$^a$</th>
<th>α-Naphthyl Acetate</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>0.10 ± 0.01</td>
<td>μmol/min/μg protein</td>
</tr>
<tr>
<td>4</td>
<td>0.22 ± 0.06</td>
<td>mean ± SD, n = 3-5</td>
</tr>
<tr>
<td>5</td>
<td>0.57 ± 0.05</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>0.84 ± 0.05</td>
<td></td>
</tr>
<tr>
<td>7.5</td>
<td>0.72 ± 0.05</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>0.30 ± 0.05</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>0.12 ± 0.05</td>
<td></td>
</tr>
</tbody>
</table>

$^a$ μmol/min/μg protein; mean ± SD, n = 3-5.

The gel mobilities of species d (128 kDa) and e (28 kDa) in the solubilized gp70 preparation were unaffected by the presence or absence of DTT (Figure 5A, lanes 3 and 4). These polypeptides remained in the unbound fraction after Con A chromatography of the solubilized gp70 preparation (Figure 5A, lanes 2 and 5) and were inactive in the esterase assay, whereas Con A-purified gp70 had esterase activity. No other polypeptides were detected in the solubilized gp70 preparation, as judged by Coomassie blue staining of 15% polyacrylamide-SDS gels (data not shown). Predictably, DTT (and β-mercaptoethanol; data not shown) inhibited the esterase activity of gp70 in a dose-dependent manner (Figure 5C), indicating that the integrity of its intramolecular disulfide bond(s) was needed for enzyme activity. Since disulfide-bonded dimers of gp70 were not observed on SDS gels or immunoblots, the reductants were breaking intramolecular disulfide bonds in the protein that were responsible for its altered SDS gel mobility. We conclude from the activity assays and SDS-PAGE analyses that monomeric gp70 was an esterase.

Growth and Development Improved by Overexpressed gp70 but Retarded in Its Absence

Before comparing growth and development parameters of strains AT-K$_{neg}$ (gp70-minus) and AT-K2 (gp70 overexpressor) to AX2, we examined the levels of gp70 in their cells and spores and assayed for secreted gp70. Indirect immunofluorescence microscopy of axenically grown, vegetative cells revealed what appeared to be a preferential distribution of esterosomes at the periphery of AX2 and AT-K2 (Figures 6B and 6D). There was an average of close to seven esterosomes (6.65 ± 3.81, mean ± SD, n = 40) per optical section from axenic cultures grown to 5 × 10$^6$ cells/ml in AX2 cells. This increased to nearly 30 esterosomes (29.02 ± 13.52, mean ± SD, n = 40) per optical section in similarly grown AT-K2 cultures (Figure 6D). Esterosomes were not observed in AT-K$_{neg}$ cells (Figure 6F). The observations of stained cells were confirmed and quantified by immunoblotting (Figure 6G). Using immunoaffinity-purified gp70 as a standard, AT-K2 cells had 4.5-fold more gp70 than AX2 cells. This was close to the 6.5-fold difference in esterase activity seen in preparations of cytoskeletons from these strains [7]. The gp70 content of axenically grown, midlog AX2 cells was estimated to be 0.06 ± 0.01% (mean ± SD; n = 4) of the total SDS-extractable protein. Although gp70 was absent (Figures 6F and 6G), lysates of axenically grown AT-K$_{neg}$ had 73% of the esterase activity (using the substrate α-naphthyl acetate) measured in AX2 lysates.

Figure 5. DTT altered the electrophoretic mobility of gp70 and inhibited its esterase activity. (A) Coomassie blue-stained 10% polyacrylamide-SDS gel of fractions, from Con A chromatography of solubilized gp70, in the presence (lanes 1–3) and absence (lanes 4–6) of 40 mM DTT. Lanes 1 and 6, α-methyl mannoside-eluted fraction containing gp70 (c, subunit of Con A); lanes 2 and 5, unbound fraction (d and e, non-Con A-binding species); lanes 3 and 4, solubilized gp70 (g, gp70) before exposure to Con A matrix. Species a and b, evident in the solubilized gp70 fraction, were separated from gp70 only in the presence of DTT (lane 3). Equal volumes of fractions were loaded. M, prestained molecular markers. (B) Immunoaffinity-purified gp70 (0.2 μg) in the presence (lanes 1 and 2) or absence (lanes 3 and 4) of 40 mM DTT. Lanes 1 and 2, unbound fraction (d and e, non-Con A-binding species); lanes 3 and 4, solubilized gp70 (g, gp70) before exposure to Con A matrix. Species a and b, evident in the solubilized gp70 fraction, were separated from gp70 only in the presence of DTT (lane 3). Equal volumes of fractions were loaded. M, prestained molecular markers. (C) The esterase activity (using the substrate α-naphthyl acetate) of solubilized gp70 was inhibited by DTT (mean ± SEM, n = 4). The x-axis indicates the amount of DTT present in the standard esterase assay using α-naphthyl acetate.
Comparison by microscopy of axenically grown cells revealed differences in the size of AT-\(K_{\text{neg}}\) cells compared to AX2 (and AT-K2) cells (Figure 8). AT-\(K_{\text{neg}}\) cells in log-phase growth were heterogeneous in size with an average cell volume 1.88-fold greater than AX2 cells. Staining with DAPI showed that the larger AT-\(K_{\text{neg}}\) cells also usually contained more nuclei compared to AX2 (Figures 8B and 8D). A small but consistent shift in roughly 35% of the population toward multinucleated cells was evident when the nuclei of AT-\(K_{\text{neg}}\) cells were counted, whereas less than 5% of AX2 cells were multinucleated (Figure 8E). The large, multinucleated AT-\(K_{\text{neg}}\) cells were found throughout log-phase growth, which supported the idea that cytokinesis was affected by the absence of gp70 rather than the nutritional state of the cells. A similar shift of cell populations toward larger numbers of nuclei also has been reported in profilin I/II-minus mutants [41]. Another observed abnormality of axenically grown AT-\(K_{\text{neg}}\) cells was the frequency of large cells containing giant vacuoles. In established log-phase cultures, up to 5% of the total cells contained the large vesicles while in wild-type AX2 cultures under the same conditions giant vacuoles are rare (1 in \(10^6–10^8\) cells) [42].

Cultures of AX2, AT-K2, and AT-\(K_{\text{neg}}\) in liquid media were monitored to determine if the absence and overexpression of gp70 affected axenic growth (Figure 9A). AT-K2 grew faster, with a generation time (GT) of 10.8 ± 1 h \((n = 4)\), than AX2 (GT = 12.2 ± 0.7 h; \(n = 4\)), and reached a maximum density of approximately 1.5 × 10^7 cells/ml. AT-\(K_{\text{neg}}\) cells, however, under the same conditions and after the same period of time, had a much longer GT of 17 ± 1.3 h \((n = 4, P = 0.0007, \text{Student’s }t\text{ test})\) and reached a maximum density only of approximately 6 × 10^6 cells/ml before cell death.

To determine if the reduced density of AT-\(K_{\text{neg}}\) was a consequence of impaired pinocytosis, the fluid-phase uptake rates of the three strains were tested using the marker FITC-dextran (Figure 9B). Pinocytosis of AT-\(K_{\text{neg}}\) cells during a 60-min period was 20% less than that of AX2 and AT-K2, which had similar rates. AT-\(K_{\text{neg}}\) cells that had reached their maximum density (6 × 10^6 cells/ml) had little to no pinocytosis activity (data not shown), which was likely why these cultures reached stationary phase at a lower density than cultures of AX2 and AT-K2.

Growth of the strains in bacterial suspensions also was monitored (Figure 10A). AT-K2 grew the fastest, and AT-\(K_{\text{neg}}\) doubled more slowly than AX2. After growth in KA suspensions, the average volume of AT-\(K_{\text{neg}}\) cells was 1.12-fold greater than that of AX2 cells. This was a significant reduction from the 1.88-fold volume difference during axenic growth. Phagocytosis assays of axenically grown AX2, AT-K2, and AT-\(K_{\text{neg}}\) revealed that AT-K2 was 22% more active than AX2, while AT-\(K_{\text{neg}}\) had only 44% the activity of AX2 (Figure 10B; mean ± SD, \(n = 7–8\)).
Growth of colonies of AX2, AT-K2, and AT-K\textsubscript{neg} on bacterial lawns followed the pattern of suspension cultures (Figure 10C). For the first week, the three cell lines had similar expansion rates. By day 8, the lines began to show differences. AX-K2 colonies expanded more quickly than AX2 and AT-K\textsubscript{neg} lagged behind AX2.

AX2, AT-K2, and AT-K\textsubscript{neg} exhibited differences in their rates of development once aggregates were formed (Table 3). All three strains took roughly 8 h to aggregate. AX2 and AT-K2 both produced mature fruiting bodies by 24 h. However, it took over 36 h for AT-K\textsubscript{neg} to reach this stage. (Student’s \textit{t} test, \(P = 0.0001\)). Another difference that reflected altered development was the number of fruiting bodies formed per square centimeter at the core of the expanding colony on bacterial lawns. The numbers for AX2, AT-K2, and AT-K\textsubscript{neg} were 34 ± 11, 43 ± 12, and 5 ± 3 fruits/cm\(^2\), respectively (\(n = 7\)). The number of fruiting bodies formed by AT-K2 was moderately higher than that of AX2 (Student’s \textit{t} test, \(P = 0.1818\)), while this number in AT-K\textsubscript{neg} cells was significantly lower than that of AX2 (Mann-Whitney test, \(P = 0.0006\)). In addition to the delayed formation of fruits, starved AT-K\textsubscript{neg} cells typically formed larger slugs, compared to those of AX2 and AT-K2 (data not shown).

**DISCUSSION**

**Function of gp70 during Vegetative Growth**

The recruitment of gp70 to phagosomes (Figures 2 and 3), presence in lysosomes (Figure 4), and esterase activity support a role for gp70 in the metabolism of endocytosed molecules in growth phase cells. The close proximity of esterosomes to the cell membrane (Figures 6B and 6D) may facilitate the fusion of esterosomes with newly formed phagosomes or endosomes or the secretion of gp70, if it is used for spore coat degradation as suggested [6]. Electron microscopy studies led to the conclusion that crystals (presumably esterosomes) disappear after spore germination [4]. However, our antibodies detected the presence of gp70 in spores, newly germinated AX2 cells (Figures 6H and 7), and growth-phase cells (Figures 1, 6B, and 6G), indicating that gp70 plays a role in axenic cell growth. Also, we detected only very low amounts of gp70 in culture media of log-phase cells, which argues for an intracellular rather than extracellular role for the enzyme during vegetative growth.

The assays of solubilized gp70 and Con A-purified gp70 established unequivocally the esterase activity ascribed to...
gp70 in a previous study [7], and further quantified its substrate range and pH dependence \textit{in vitro} (Tables 1 and 2). Gp70 was active in a pH range from pH 5 to pH 7.5, which would permit it to function in both acidic lysosomes [36–38] and neutral postlysosomal compartments [39]. Its catalytic activity, like other lysosomal enzymes in \textit{D. discoideum} [43], was enhanced at pH 6, close to the measured pH of \textit{D. discoideum} lysosomes [36, 38]. Since both phagosomes and pinosomes are acidified by the vacuolar H\(^{+}\)-ATPase (Figure 3) [34, 37, 44–46], the more effective activity of gp70 at lower pH is consistent with our suggested role of gp70 in the hydrolysis of endocytosed nutrients. Possibly the lower pH also plays a role in the transition of gp70 from a crystalline state (Figure 1; estersomes) to a soluble enzyme when fused with phagosomes and pinosomes.

![Figure 8](image)

**Figure 8.** Absence of gp70 hindered cytokinesis in axenically grown AT-K\(_{\text{neg}}\) cells. Bright-field images (A, C) and DAPI staining (B, D) showed that AT-K\(_{\text{neg}}\) cells (C, D) were larger in size and multinucleated than those of AX2 (A, B). Bar = 20 µm. (E) Number of nuclei per cell of axenically grown AX2 (stippled) and AT-K\(_{\text{neg}}\) (solid).

![Figure 9](image)

**Figure 9.** Axenic growth and fluid-phase endocytosis of AT-K\(_{\text{neg}}\) were reduced compared to AX2 and AT-K2. (A) Cell densities of axenically grown cultures of AX2 (diamonds), AT-K2 (circles), and AT-K\(_{\text{neg}}\) (squares) were monitored (mean ± SD, \(n = 4\)). (B) Pinocytosis was measured using the fluid-phase marker FITC-dextran (mean ± SD, \(n = 5–7\)).

Nonspecific esterase activities have been reported in \textit{D. discoideum} [47], as were esterase activities that specifically cleave long chain (lauric, myristic, and palmitic) fatty acids from the bioactive, hydrophobic lipid A moiety of lipopolysaccharides (LPS) [48, 49]. Cleavage of another acyl chain, the amide-linked hy-droxymyristic acid of lipid A, requires amidase activity [50, 51], which was not detected with solubilized gp70. Extracts from other organisms (the slime mold \textit{Physarum polycephalum} [52] and protozoan \textit{Acanthamoeba} [53]) that consume bacteria presumably contain esterases because they are able to deacylate LPS. To our knowledge, except for an acyloxyacyl hydrolase from human neutrophils that deacylates LPS [54], enzymes responsible for metabolizing LPS, particularly the lipid A portion, have not been identified and examined. Because of the interest in how animals respond to and neutralize gram-negative bacteria, future studies will test if LPS is a substrate for gp70. However, since the absence of gp70 had a modest effect on \textit{D. discoideum} growth on bacteria (Figure 10), other esterases, including the related D2 protein, must be involved in the deacylation LPS in this organism.
Esterase gp70 and Growth and Development of *Dictyostelium discoideum*

AT-K2 led to modestly superior growth and phagocytic activity (Figures 9 and 10) and may have been due to the presence of low levels of gp70 acting to deacylate lipid molecules. These types of benign consequences are observed frequently with other overexpression mutants of *D. discoideum* [55–57]. Assuming that gp70 was an active esterase in axenically grown cells, we postulate that its absence delayed the digestion of pinocytosed nutrients or caused harmful compounds to accrue. Consequently, the endosomal network was mildly disrupted, leading to impaired cell division. We attribute the longer doubling time of AT-K$_{neg}$ cells and their low culture densities (Figure 9A) to irregularities in the endosomal membrane network and reduced pinocytosis (Figure 9B).

**Role of gp70 in Development**

The absence of gp70 was expected to affect development (Table 3), since gp70 is developmentally regulated [6]. The slower development and smaller number of fruiting bodies from AT-K$_{neg}$ on bacterial lawns explain the slower expansion of these colonies, compared to AT-K2 and AX2, that became more pronounced with time (Figure 10C). Also developmentally regulated is the protein D2, presumed to be an esterase, which is found in esterosomes with gp70 and related to it by 52% amino acid sequence identity [6, 58]. When starved, axenically grown antisense mutants of D2 are delayed in streaming and aggregate formation [58]. Since both gp70 and D2 are expressed during development, the ability of AT-K$_{neg}$ cells to form eventually fruiting bodies may be due to the substitute activity of D2. It would be interesting to test if a double knockout of the gp70 and D2 genes has an additive effect on inhibiting development. Alternatively, other esterases [47] may compensate for gp70, particularly during growth on bacteria when its absence had minimal effects. The idea that there are redundant lysosomal enzymes is supported by the recent report that the absence of the *D. discoideum* cathepsin D protease has the sole effect of slightly delaying development while other cell activities are normal [59].

In conclusion, gp70 was an active esterase present in digestive/endocytic vesicles of vegetative *D. discoideum* amoebae. Immunolocalization and biochemical analyses supported a role for gp70 in the deacylation of fatty acids from lipids such as bacterial LPS. Comparative physiology studies of strains that either overexpress or lack gp70

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**Figure 10.** Bacterial growth and phagocytosis of AT-K$_{neg}$ were reduced compared to AX2 and AT-K2. (A) Densities of AX2 (diamonds), AT-K2 (circles), and AT-K$_{neg}$ (squares) cultures grown in bacterial suspensions (mean ± SD, n = 4). (B) Phagocytosis was measured using TRITC-yeast (mean ± SD, n = 7–8). (C) Diameters of colonies grown on KA lawns (mean ± SD, n = 4).

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**Table 3**

Time Elapsed for Development$^a$

<table>
<thead>
<tr>
<th></th>
<th>Aggregates</th>
<th>Fruiting bodies</th>
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<tbody>
<tr>
<td>AX2</td>
<td>7.5 ± 0.5</td>
<td>23.3 ± 1.5</td>
</tr>
<tr>
<td>AT-K2</td>
<td>7.9 ± 0.7</td>
<td>23.7 ± 1.3</td>
</tr>
<tr>
<td>AT-K$_{neg}$</td>
<td>7.9 ± 0.9</td>
<td>36.9 ± 2.4</td>
</tr>
</tbody>
</table>

$^a$Hours; mean ± SD, n = 7-11.

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Although AT-K$_{neg}$ (gp70-minus) cells had measurable esterase activity, gp70 was important for growth because its absence disrupted cytokinesis, producing large cells (Figure 8), and reduced phagocytic activity, leading to slower growth on bacteria (Figure 10). In contrast, overexpression of gp70 in
provided evidence that the enzymatic activity of gp70 was needed for normal growth and development. Future studies will examine the mechanism of recruitment of gp70 to endosomes and regulation of its enzymatic activity.

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References


