1-1-2001

Giant Vacuoles Observed in *Dictyostelium discoideum*

Aidong Yuan  
*University of Nebraska - Lincoln*, dyuan@unlserve.unl.edu

Catherine P. Chia  
*University of Nebraska - Lincoln*, cchia1@unl.edu

Follow this and additional works at: [http://digitalcommons.unl.edu/bioscimicro](http://digitalcommons.unl.edu/bioscimicro)

[http://digitalcommons.unl.edu/bioscimicro/80](http://digitalcommons.unl.edu/bioscimicro/80)
Introduction

Fluid-phase endocytosis, pinocytosis, is the ingestion of fluid and solutes via small uniform vesicles called pinosomes (≤ 150 nm in diameter (Alberts et al., 1994). Pinosomes larger than 0.2 μm are called macropinosomes in mammalian cells (Swanson and Watts, 1995), and the same term has been applied to large endocytic vesicles in amoebae of axenically-grown strain AX2 (0.6 μm in diameter (Thilo and Vogel, 1980) and that of macropinosomes, 1.3–1.9 μm (Hacker et al., 1997). In D. discoideum, the average diameter of primary pinosomes is about 0.6 μm (Thilo and Vogel, 1980) and that of macropinosomes, 1.3–1.9 μm (Hacker et al., 1997). In the course of microscopic studies on endocytosis, we observed “giant” vacuoles, which were >4 μm in diameter, with some having diameters of up to 30 μm which varied in their frequency among different strains. Because the vacuoles took up a fluid-phase marker, fluorescein-labeled dextran, they appeared to be part of the endolysosomal system, and could reflect a delay or interruption of endosome processing.

Materials And Methods

Cell growth

Dictyostelium discoideum cells (AX2, KAX3, and transformed cell line AT-Kneg, a strain that lacks gp70, a 70 kDa glycoprotein with esterase activity; Chia et al., 1998) were grown typically in shaken cultures to 1 to 6 × 10⁶ cells/ml at 20°C in 50 ml of HL5 nutrient media (Watts and Ashworth, 1970). The inclusion of G418 (10 μg/ml), needed to maintain the transformant, did not promote the formation of giant vacuoles in the parental AX2 cells. Culture densities were monitored using a haemocytometer to count cells.

Microscopy

For viewing intracellular giant vacuoles, log-phase cells were withdrawn from cultures, applied to a haemocytometer and examined immediately. When cells were on the haemocytometer for 1 to 3 h, protruding giant vacuoles were observed. The percent of cells with giant vacuoles (with diameters ≥4 μm) was determined by
counting cells in the 1 mm² grids of the haemocytometer. Typically, nine grids were counted for each of five to ten cultures of each strain. Cell viability was monitored using 0.4% trypan blue (Ahern, 1992). Cells were observed with a Nikon Labophot microscope (Melville, NY, USA) using either bright field (40× objective, NA 0.65) or epifluorescence (using a 60× Plan Apochromat oil objective, NA 1.4) optics. Images were acquired with a 3-chip color CCD camera (DC-330, DAGE-MTI, Inc., Michigan City, IN, USA) and a Scion CG-7 RGB color frame grabber (Scion Corp., Frederick, MD, USA) using Scion Image (an extended version of NIH Image available at http://www.scioncorp.com) and imported into Photoshop 5.0™. Figure 3 was obtained with confocal microscopy model MRC-1024ES (Bio-Rad Laboratories, Hercules, CA, USA) at the University of Nebraska Center for Biotechnology Microscopy Core Facility. For some experiments, fluorescein 5-isothiocyanate (FITC)-dextran (Sigma Chemical Co., St Louis, MO, USA) was added to the HL5 to a final concentration of 2 mg/ml (Vogel, 1987). Cells were withdrawn at various times up to 10 h, and collected by centrifugation at 200× g at 4°C and washed three times with Sorensen’s buffer (14.6 HIM KH₂PO₄, 2 mM Na₂HPO₄, pH 6.1) before observation and fixation.

For immunofluorescence microscopy, 50 μl of washed cells, resuspended at 5 × 10⁵ cells/ml in cold Sorensen’s buffer, were applied to coverslips. Settled cells were fixed with 3.7% formaldehyde in 17 mM sodium phosphate buffer, pH 6.8, for 15 min at 20°C, and permeabilized with chilled methanol (–20°C) containing 1% formaldehyde for 5 min (Fukui et al., 1987). After blocking with 1% BSA in PBS (phosphate-buffered saline; 150 mM NaCl, 10 mM sodium phosphate, pH 7.4) for a minimum of 15 min at room temperature, slides were incubated for 1 h at room temperature with the culture supernatant of hybridoma N2 (1:10 in 1% BSA in PBS) specific for the 100 kDa subunit of the vacuolar H⁺-ATPase of D. discoideum (V-H⁺-ATPase) (Fok et al., 1993). Coverslips were washed three times with PBS containing 0.05% Tween 20, and cells were stained for 1 h with Rhodamine Red-X™-goat anti-mouse IgG in 1% BSA in PBS (1:150; Jackson ImmunoResearch Laboratories, Inc., West Grove, PA, USA). Samples incubated with secondary antibody when primary antibody was omitted had no significant fluorescent signals. Coverslips were mounted in buffered polyvinyl alcohol containing 50 mg/ml 1,4-diazobicyclo-(2,2,2)-octane (Aldrich Chemical Co., Milwaukee, WI, USA) (Fukui et al., 1987).

Results

Intracellular giant vacuoles in axenically grown AX2 cells

Intracellular giant vacuoles, >4 μm in diameter, were rare in axenically grown AX2 amoebae (about 1 in 10⁶–10⁸ cells). Typically, there was one giant vacuole, round or oval-shaped, per cell (Figure 1A and 1B). The giant vacuoles were dynamic structures since they would appear and disappear over a 30 min period (data not shown). Cells with the giant vacuole(s) were usually, though not always, up to 50–60 μm in diameter (Figure 1A and 1B). Nuclear staining with 4, 6-diamino-2-phenylindole of these large cells containing intracellular giant vacuole(s) revealed multiple nuclei (>10 to 20 per cell; data not shown).

Protruding giant vacuoles produced under hyperosmotic conditions

When monitored over several hours in a haemocytometer, over 5% of the AX2 cells left in HL5 formed giant vacuoles that bulged noticeably from the cells. A similar frequency of protruding giant vacuoles was observed in all strains examined. The vacuoles extended from either an individual cell (Figure 1C) or a cell in an aggregate (Figure 1D), and often had dimensions equivalent to or larger than the cell (Figure 1C and 1D). As seen with the intracellular vacuoles, the protruded vacuoles were dynamic. They would retract gradually into the cell during a 20-30 min interval (Figure 2). Confocal microscopy established that a vacuole was indeed retracting and not the rotation of the cell that gave the illusion of a shrinking vacuole. Figure 3 shows images collected from an AX2 cell (similar to the one shown in Figure 2, panel D) indicating that the cell and retracting vacuole remained in the same optical plane. Thus the decreased size of the vacuole was due to its retraction into the cell rather than a change in its position relative to the viewer.

Fluid-phase marker FITC-dextran accumulated in giant vacuoles

To determine if the giant vacuoles were derived from endosomes, the fluid-phase marker FITC-dextran was added to log-phase cells growing in nutrient media. Small, fluorescent pinosomes (<1 μm in diameter) were seen in cells within 30 min to 1 h, consistent with the high pinocytic activity of the cells. Extended culture...
for 4 h revealed larger fluorescent pinosomes (about 2 μm in diameter; data not shown), suggesting the fusion of small pinosomes into larger ones. After lengthier growth periods in the presence of FITC-dextran, giant vacuoles containing the marker were both inside (Figure 4A and 4B) and protruding from a cell (Figure 4C and 4D). Inclusion of FITC-dextran in HL5 did not increase the frequency of giant vacuoles in all strains examined. To ensure that cells with protruding giant vacuoles were viable, trypan blue was added to monitored samples. For up to 5 h, those cells left on the haemocytometer and with protruding vacuoles excluded trypan blue, indicating their continued viability.

**Vacuolar H⁺-ATPase enrichment in the membranes of intracellular giant vacuoles**

Further confirmation that the giant vacuoles were related to endosomes was the presence of a H⁺-ATPase in...
the vacuolar membranes. The monoclonal antibody N2 recognizes the 100 kDa subunit of the *D. discoideum* vacuolar H\(^+\)-ATPase, which is localized to endomembrane structures (Nolta et al., 1994), and the membranes of the contractile vacuole (CV) (Fok et al., 1993) in log-phase *D. discoideum* cells. Cells with giant vacuoles loaded with pinocytosed FITC-dextran and stained with N2 showed strong co-localization of the two signals (Figure 5B and 5C), indicating that the vacuoles were endocytic compartments.

**Figure 2.** Protrusion and retraction of a giant vacuole. (A-F) AX2 cell with a protruded giant vacuole was monitored and images were captured at the indicated times. The giant vacuole contracted gradually and appeared to retract into the cell. Scale bar, 10 μm.

**Figure 3.** Confocal images of an AX2 cell with a protruding giant vacuole. Panels D1 through D11 show 1 μm thick images, with a 0.3 μ overlap, from the top (D1) to the bottom (D11) of a cell similar to the one shown in Figure 2D. An asterisk labels the vacuole. Scale bar, 10 μm.
Giant Vacuoles Observed in *Dictyostelium discoideum*

Although relatively rare in AX2 cells, giant vacuoles were more common in newly germinated KAX3 cells. They occurred at an average frequency of 0.55% (SD = 0.062%, *n* = 5) of the population. Usually, there was one giant vacuole per cell (Figure 6A and 6B), but occasionally cells had two or more of these structures (Figure 6C and 6D). When left on a haemocytometer for 1-3 h, protruding vacuoles were seen also in KAX3 cells (Figure 6E). The giant vacuoles were present in only one of the cells (Figure 6F).

Discussion

The observed giant vesicles (>4 μm diameter; Figures 1, 6, & 8) were substantially larger than the pinosomes (0.6 μm diameter) (Thilo and Vogel, 1980), macropinosomes (1.3-1.9 μm diameter) (Hacker et al., 1997) and prelysosomal acidic vacuoles (<2 μm in diameter) (Padh et al., 1989) previously described for *D. discoideum*. The formation of the giant vacuoles could occur by fusion of small pinosomes (endosomes) or enlargement of the individual vesicles. The fusion model is supported by our observation that pinosomes were fewer and larger with the extended culture of cells in nutrient media containing FITC-dextran, and a study using *D. discoideum* extracts that demonstrated *in vitro* endosome-endosome fusion (Lenhard et al., 1992).
The occurrence of intracellular giant vacuoles was infrequent, and presumably due to a metabolic aberration occurring in AX2 cells during vegetative growth. It is unlikely that these structures were a consequence of flaws in the culture system because only a few cells had intracellular giant vacuoles and the vast majority appeared normal in the standard HL5 nutrient media. In addition, the frequency of giant vacuoles varied with different strains grown under the same conditions with the same media. One strain, AT-K2, which overexpresses gp70 (Chia et al., 1998) had a frequency of giant vacuole formation even lower than that of AX2 (data not shown). It remains puzzling why newly geminated KAX3 cells had a much higher frequency of intracellular giant vacuoles (Figures 6 & 7) when compared to that of cultures maintained over 1–2 months.

Figure 6. KAX3 cells had giant vacuoles. (A, B) Intracellular giant vacuoles were nearly spherical and (C, D) occasionally several were in a single cell. (E, F) Vacuoles protruding from cells are labeled with asterisks. Scale bar 10 μm.
The presence of the H\textsuperscript{+}-ATPase in some of the giant vacuoles (Figure 5) was consistent with the genesis of these structures from endocytic vesicles, which in *D. discoideum* are acidified by a vacuolar H\textsuperscript{+}-ATPase (Cardelli et al., 1989; Nolta et al., 1994; Padh et al., 1989, 1991a; Temesvari et al., 1996). Our suggestion that giant vacuoles were derived from endosomes agrees with an earlier finding that inhibition of the vacuolar H\textsuperscript{+}-ATPase

*Figure 7.* Giant vacuoles in KAX3 harbored FITC-dextran. (A, C, and E) Bright field images of giant vacuoles (indicated with arrows) inside (A, C) or protruding out from (E) cells contained the fluid-phase marker FITC-dextran (B, D, and F). Scale bar, 10 μm.
induces the formation of large intracellular vesicles in *D. discoideum* (up to 5 μm in diameter) that accumulate FITC-dextran (Temesvari et al., 1996). Possibly faulty or hindered acidification of endosomal vesicles interrupted normal processing and led to the formation of the giant vacuoles, since acidification is a condition for proper targeting of a lysosomal enzyme (Cardelli et al., 1989; Temesvari et al., 1996). Similarly, the relatively high frequency of giant vacuoles in AT-K<sub>neg</sub> cells (Figure 8) could be the result of altered endolysosome processing in the absence of the enzyme that is normally present in lysosomes (Yuan and Chia, in preparation).

An alternate explanation for the origin of giant vacuoles arises from the similarity of the protruding structures to previously described bulging vacuoles, known to contain the CV marker, alkaline phosphatase (Quiviger et al., 1978). The giant vacuoles may be part of the CV since they apparently occurred in response to the changing osmolarity of the nutrient medium. Since excretion of the vacuolar contents presumably would occur through the CV complex, the giant vacuoles may be composite structures related to both endosomes and the CV, which are evidently different organelles (Gabriel et al., 1999) but share some common molecules (e.g., the vacuolar-H<sup>+</sup>-ATPase [Heuser et al., 1993; Nolta et al., 1993] and a Rab4-like GTPase [Bush et al., 1994]). These two organelle systems may be connected under certain osmotic conditions (Padh, 1995; Padh et al., 1991a,b; Steck et al., 1997).

In summary, we report the presence of intracellular giant vacuoles (>4 μm in diameter) that occurred at varying frequencies in strains of axenically grown *D. discoideum*. Mild osmotic stress induced the formation of giant vacuoles that protruded from cells. Some intracellular vacuoles harbored the fluid-phase marker FITC-dextran and contained a vacuolar-H<sup>+</sup>-ATPase in their membranes, suggesting their derivation from endocytic vesicles. Both intracellular and protruded vacuoles harbored the fluid-phase marker FITC-dextran, implicating a relationship between the CV and endolysosomal networks.

Acknowledgments

We thank Dr A. Fok (Pacific Biomedical Research Center, University of Hawaii) for providing the N2 hybridoma supernatant, Dr C.-H. Siu (University of Toronto) for strain KAX3, and Dr Y. Zhou for the confocal microscopy images. These studies were supported by funds from the Center for Biotechnology, University of Nebraska-Lincoln and NSF Grant MCB-95133628 to C.P.C.
References


