Cyanophycin Granule Size Variation in *Aphanocapsa*

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Techniques were developed for the rapid and simple electron microscopic visualization of isolated cyanophycin granules to allow analysis of their size from cells grown under different conditions. Granules were purified by differential and renografin density-gradient centrifugation from cells grown for various times in chloramphenicol-containing media, or in nitrogen-limited media to which nitrogen was then added back, or under light limitation. Granules were then observed by electron microscopy after staining with uranyl acetate, and random granule diameters were measured. Granule size increased significantly with time following chloramphenicol treatment and following light limitation. This suggests that much of the increase in amount of cyanophycin in cells treated with chloramphenicol or light limitation is caused by increase in granule size rather than in granule number. Insignificant changes in granule size were observed during nitrogen repletion, suggesting increase in granule number, not size.

INTRODUCTION

Cyanophycin granule polypeptide [CGP, multi-L-arginyl poly(L-aspartic acid)] is a nitrogen storage protein found uniquely in cyanobacteria (Allen & Hutchison, 1980; Lawry & Simon, 1982). It is a polymer, synthesized non-ribosomally, consisting of an equal molar ratio of arginine and aspartic acid (Simon, 1973b). Formation of this compound, which is usually found in a granular form (cyanophycin granules), is induced by starvation of nutrients other than nitrogen (Allen, 1980) and by addition of a number of growth-inhibitory agents such as chloramphenicol (CM) (Simon, 1973a; Lawry & Simon, 1982).

The only published fine-structural studies of cyanophycin granules have been of thin-sectioned material (Lang et al., 1972; Allen & Weathers, 1980; Lawry & Simon, 1982) or of freeze-fractured material (Allen & Weathers, 1980). Little information is available on the mechanism of CGP synthesis such as peptide initiation, elongation and how and when granules increase in size. Allen & Weathers (1980) showed small granules in thin sections of light-limited *Aphanocapsa* in late-exponential growth, whereas larger granules were observed in stationary-phase cells. In the present work, electron microscopic techniques were developed for rapid and easy observation of granule size variation. Granules isolated from *Aphanocapsa* 6308 at various times after CM treatment or after light limitation were then analysed since the amount of CGP increases rapidly under these conditions (Allen et al., 1980). Cyanophycin granules from nitrogen-limited and regreening cells were also observed for changes in size or appearance.

METHODS

Organism and growth. *Aphanocapsa* 6308 (*Synechocystis* ATCC 27150) was routinely grown with 5% (v/v) CO₂ in air in liquid medium no. 11 (Allen, 1968) as described previously (Allen & Hutchison, 1980). CM (5 µg ml⁻¹) was added in some experiments. In other experiments cells were grown in carboys containing 121 of medium no. 11

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 Abbreviations: CGP, cyanophycin granule polypeptide; CM, chloramphenicol.
Fig. 1. Uranyl-acetate-stained purified cyanophycin granules isolated from Aphanocapsa grown under various conditions: (a) from exponentially growing cells; (b) from cells treated with CM for 12 h; (c) from cells treated with CM for 48 h. Bars, 1 μm.

with 1/40 the normal concentration of NaNO₃. After cells were starved for nitrogen, NaNO₃ was added and samples were collected at various times, during regreening for granule isolation.

**Cyanophycin granule isolation.** At various times after treatments, cells were collected, disrupted using a French pressure cell (138 MPa pressure) and cyanophycin granules were purified. The 27000 g pellet, after a 15 min centrifugation of the crude extract, was washed with distilled water followed by 2% (v/v) Triton X-100 before being resuspended in water and placed on 70-100% (v/v) renografin step gradients (Allen & Weathers, 1980). After 2 h at 18000 r.p.m. in a SW41 rotor, granule bands were removed, washed twice with distilled water and prepared for electron microscopy. Paper chromatography, after hydrolysis of this material (Allen & Weathers, 1980), showed only arginine and aspartic acid to be present; quantitative analysis of the granule bands was done by the method of Simon (1973a).

**Electron microscopy.** Washed samples (5 μl) from renografin gradients were placed on 300-mesh copper grids for 3 min. Excess liquid was removed and the grids were stained for 30 s with 1% (w/v) uranyl acetate. The grids were observed with a Zeiss 9s-2 electron microscope operated at 60 kV. Random micrographs were taken and granule diameters were measured from enlarged prints. Significance of the results was determined using Duncan's test (Bruning & Kintz, 1968).

**RESULTS**

Purified cyanophycin granules were observed from cells harvested before (0 h) and 6, 12, 24 and 48 h after addition of CM. Fig. 1(a–c) shows representative fields of granules from these various sampling times. There was a marked increase in granule size with time after CM addition. Quantitative data are shown in Fig. 2. Mean granule diameter ranged from 181 nm (isolated from exponentially growing cells) to 446 nm (isolated from cells treated for 48 h with CM). Lack of overlap of the standard errors of the means shows that the differences observed in granule diameter are greater than any error that could result from inaccuracy in measurement between groups. Standard deviation of the mean values for each sampling showed a slight increase with time (data not shown), but the largest of the granules isolated from cells before CM treatment were smaller than the smallest of those 48 h after the addition of CM. Note the amorphous material associated with the granules isolated from exponentially growing cells (Fig. 1a).

Granules isolated from nitrogen-limited or regreening nitrogen-starved cells were difficult to purify. As observed for granules from exponentially growing cells (Fig. 1a), much amorphous material was associated with the granules throughout the purification process. The preparations were green throughout isolation. Fig. 3(a) shows purified material from cells starved for nitrogen for 98 h (0·3% CGP on a dry weight basis). Fig. 3(b) shows granules purified from cells harvested 15 h after NaNO₃ was added (3·4% CGP on a dry weight basis).

Purified granules from cells grown to late-stationary phase (11 d growth) had a mean diameter of 1390 nm (245 randomly selected granules were measured), three times that of granules isolated from cells grown in CM for 48 h. Fig. 3(c) shows an example of granules isolated from...
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Fig. 2. Mean diameter of *Aphanocapsa* cyanophycin granules versus time after CM addition. Standard errors of the means are shown; *n* = number of granules counted.

Fig. 3. Uranyl-acetate-stained purified cyanophycin granules from *Aphanocapsa*: (a) from cells starved for nitrogen for 98 h; (b) from nitrogen-starved cells 15 h after NaN03 was added; (c) from light-limited (stationary-phase) cells. Bars 1 μm.

these light-limited cells. Much more variation in size was observed than in the granule populations isolated from CM-treated cells.

**DISCUSSION**

The present results indicate that increase in cyanophycin granule polypeptide in CM-treated and stationary-phase *Aphanocapsa* appears to be by an increase in granule size and not by an increase in *de novo* synthesis of granules.

When cells are growing exponentially under favourable conditions, CGP is present in low amounts, only 0.1% of the dry weight of the cell in *Anabaena cylindrica* (Simon, 1973a, b) or 1.5% in *Aphanocapsa* (Allen et al., 1980). When cells of *Aphanocapsa* were in light-limited stationary phase, CGP increased to 16% of the dry weight; cells treated with CM for 48 h accumulated CGP to 11% of their dry weight (Allen et al., 1980). Light-limited Calothrix *marchica*, observed in thin section, contained cyanophycin granules up to 740 nm in diameter (Tahmida Khan & Godward, 1978), and analysis of thin sections of light-limited *Aphanocapsa* suggests that granules reach a diameter of 1900 nm (M. M. Allen, unpublished). This compares well with the mean of 1390 nm calculated in the present work. If the granules are assumed to be spherical, an increase in diameter from 181 nm to 446 nm 48 h after CM addition suggests approximately a 14-fold increase in volume. This amount compares well with the quantity of CGP shown to be present in CM-treated *Aphanocapsa* by chemical analysis (Allen et al., 1980).
An immediate linear increase in CGP (measured by analysis of arginine content), followed by a decrease in amount, is seen in nitrogen-starved cells to which nitrate is added (Allen & Hutchison, 1980). That no significant differences could be observed, over 30 h, in the size of granules from such cells, although the amount of CGP increased tenfold, at 15 h after regreening began, suggests that new granules were formed under these conditions.

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