An Ultrastructural Investigation into the Role of Calcium in Oosphere-initial Development in *Saprolegnia diclina*

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Prior to formation of oosphere initials in septum-delimited oogonia of *Saprolegnia diclina*, the volume density (volume fraction) of central vacuoles containing characteristic clusters of electron-opaque, granular material increased from 6.0 to 54.2% whereas dense-body vesicles decreased in volume density from 30.2 to 9.0%. During this time, dense-body vesicles were reduced in number by 50%, their mean diameter decreased from 0.861 to 0.697 μm and the percentage of vesicles with more than one dense-body granule increased from 5.4 to 20.2. The volume density of lipid bodies did not change, whilst that of the cytoplasmic matrix decreased from 40.0 to 18.2%. Volume densities of mitochondria and nuclei remained constant relative to the peripheral oogonial protoplasm until central vacuoles had enlarged, but then decreased as oosphere initials formed. Small cytoplasmic vesicles were associated both with Golgi dictyosomes and with dense-body vesicles having blebbed membranes. Similar small vesicles were present close to both central vacuole membranes and oogonial walls.

Oogonia grown in calcium-deficient medium showed a high level of oogonium and oosphere abortion. Aborted oogonia showed extreme protoplast disorganization. In unaborted oogonia grown under calcium-deficient conditions, central vacuoles contained crystalline inclusions, and some oogonia did not form secondary wall layers. Organelles were of normal appearance but calcium deficiency increased the volume densities of mitochondria and nuclei at early stages of oogonial development and of dense-body vesicles and peripheral vacuoles at later stages; the volume density of lipid bodies was reduced at all stages. The numerical density of dense-body vesicles progressively increased during oogonial development and was greater at all developmental stages than in oogonia grown under conditions of calcium sufficiency. At late stages of oogonial development, calcium deficiency reduced both the mean diameter of dense-body vesicles and the fraction of them having two or more dense-body granules.

**INTRODUCTION**

Calcium ions may be intimately involved in control at the cellular level (Bygrave, 1978) and can influence reproduction in Oomycetes and other fungi (Bollard & Butler, 1966; Lenny & Klemmer, 1966; Elliott, 1972). McCann & Stuart (1973) found Ca²⁺ to be essential for growth of *Saprolegnia diclina*. They also showed that deficiency of Ca²⁺ in the culture medium specifically affected the normal cleavage into oospheres of the protoplasm of developing oogonia, resulting in oogonial abortion. In the related genus *Achlya*, Ca²⁺ deficiency prevents zoospore formation (Griffin, 1966) and inhibits amino acid uptake and protein synthesis (Cameron & LeJohn, 1972).

Formation of oosphere initials in oogonia of members of the Saprolegniaceae, including *S. diclina*, is preceded by formation of a large central vacuole or vacuole system without

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change in oogonial size (DeBary, 1881; Humphrey, 1893; Hartog, 1895; Trow, 1899; Fletcher, 1978). Gay et al. (1971) suggested that the central vacuole system in oogonia of *Saprolegnia ferax* was formed by enlargement and coalescence of specialized vesicles (dense-body vesicles) and showed an overall increase in vesiculation during oogonial development, but they did not distinguish in their quantitative study between dense-body vesicles and central vacuoles or investigate quantitative changes in other organelles. DeBary (1881), Hartog (1895) and Trow (1899) described formation of oosphere initials in several members of the Saprolegniaceae as involving the progressive drawing into heaps of a thin, peripheral layer of protoplasm surrounding an already fully-enlarged central vacuole. In contrast, Howard & Moore (1970) and Gay et al. (1971) suggested, after ultrastructural studies, that oosphere initials are cut out by cleavage vacuoles that develop centrifugally while central vacuoles are still enlarging.

This paper presents qualitative and quantitative observations on the ultrastructure of developing oogonia of *S. diclina* that help to elucidate further the changes within oogonia that lead to the formation of oosphere initials. Observations on changes in oogonial ultrastructure induced by growth under conditions of Ca\(^{2+}\) deficiency are also included.

**METHODS**

*Organism and culture.* Stock cultures of *Saprolegnia diclina* Humphrey were maintained on a medium containing (g l\(^{-1}\) in deionized water): glucose 1·66; sodium glutamate 0·066; DL-methionine 0·016; KH\(_{2}\)PO\(_{4}\), 0·046; K\(_{2}\)HPO\(_{4}\), 0·056; MgCl\(_{2}\), 6H\(_{2}\)O, 0·33; MnCl\(_{2}\), 4H\(_{2}\)O, 0·02; CaCl\(_{2}\), 6H\(_{2}\)O, 0·006; FeCl\(_{3}\), 6H\(_{2}\)O, 0·0004; ZnSO\(_{4}\).7H\(_{2}\)O, 0·01; and solidified with 2 % (w/v) agar (Oxoid no. 1). Cultures for electron microscopy were grown for 5 to 10 d at 20 °C in unsolidified, quarter-strength medium containing 6·85 µM-CaCl\(_{2}\) (medium used by McCann & Stuart, 1973; M. McCann, personal communication) in glass Petri dishes inoculated with plugs from stock cultures (high Ca\(^{2+}\) regime), or in medium containing 1·71 µM-CaCl\(_{2}\) for 5 d then transferred to CaCl\(_{2}\)-free medium for 5 to 10 d (Ca\(^{2+}\)-free regime).

*Electron microscopy.* Portions of oogonia-bearing mycelium were fixed for 20 min in a 2 % (w/v) formaldehyde/2 % (v/v) glutaraldehyde mixture, rinsed three times with water, transferred to 1 % (w/v) osmium tetroxide for 1 h then rinsed three times with water, all at room temperature and buffered at pH 7·2 with 0·05 M-sodium cacodylate. Fixed material was dehydrated in an acetone/water series with retention for 2 to 4 h in 50 % (v/v) acetone saturated with uranyl acetate, and embedded in Araldite or Epon. Sections were stained at room temperature with 5 % (w/v) aqueous uranyl acetate for 30 min followed by lead citrate for 15 min.

*Stereology.* Oogonia for stereological sampling were selected by light microscopy after fixation and embedding. Sampling was carried out on (i) oogonia which were already septum-delimited but in which the central vacuole was still small (vacuole diameter about 25 % of the total oogonium diameter; young oogonia), (ii) septum-delimited oogonia in which the central vacuole was large (vacuole diameter about 80 % of the total oogonium diameter; late pre-oosphere initial stage oogonia) and (iii) oogonia in which oosphere initials had already formed (oosphere initial stage oogonia). Exact measurements of total oogonium and central vacuole diameters of young and of late pre-oosphere initial stage oogonia were made using a light microscope with a ×10 objective after trimming of blocks but before sectioning, and the fractions of the total oogonium volume occupied by central vacuoles and peripheral protoplasm, respectively, were calculated for each oogonium used. Percentage volume densities of selected organelles (volume fraction of organelle in containing protoplasm × 100) were determined on electron micrograph prints at ×10000 magnification from at least five oogonia from each Ca\(^{2+}\) regime at each developmental stage by point sampling methods (Weibel, 1973) using a 1 cm square sampling lattice, with sampling restricted to the peripheral protoplasm only. Sufficient micrographs were sampled to provide a total test point number of at least 6000 for each Ca\(^{2+}\) regime at each developmental stage. Mean diameters and numerical densities (number per unit volume of containing protoplasm) of the dense-body vesicle component were determined using the method described by Weibel (1973) with any non-circular profiles treated as circles of equal area. Volume density and numerical density values relative to the peripheral protoplasm in high Ca\(^{2+}\) regime young and late pre-oosphere initial stage oogonia were converted to values relative to the total oogonium volume (peripheral protoplasm plus central vacuoles) by multiplication by the fraction of the oogonium volume occupied by peripheral protoplasm. This conversion was made individually for each oogonium sampled. The significance of results was assessed using standard two-sample significance tests as described by Cooper (1969). Pairs of numerical density values for the same organelle at consecutive stages of development having a difference of low statistical significance (P > 0·4) were combined to provide a common estimate.
RESULTS

High Ca²⁺ regime

In young, septum-delimited oogonia (Fig. 1), central vacuoles contained characteristic clusters of granular material (Fig. 2). Electron-opaque granules, similar in distribution to the mottled granules described by Gay et al. (1971), were present in vacuoles of some oogonia (Fig. 10) but were not found consistently. In addition to nuclei, mitochondria, lipid bodies, dense-body vesicles, and Golgi dictyosomes with associated small vesicles (Figs 2, 4), the peripheral protoplasm of young oogonia contained small vacuoles (peripheral vacuoles) with clusters of granular material similar to those in central vacuoles (Fig. 3). Serial sections showed that most of the vacuole or vesicle-like profiles within the peripheral protoplasm, except for peripheral vacuoles and small vesicles similar to those associated with Golgi dictyosomes, contained dense-body granules (Figs 5, 6, 7). These granules were surrounded by an often irregular layer of material of greater electron opacity (Figs 3, 8, 9, 11). Some profiles of dense-body vesicles contained two dense-body granules either free (Fig. 8) or joined by electron-dense material (Fig. 9). Some dense-body vesicles showed blebbing and were associated with clusters of small vesicles (Fig. 11). In late pre-oosphere initial stage oogonia (Fig. 12) the large central vacuoles contained scattered membrane profiles and clusters of electron-opaque granular material (Fig. 15) similar to the clusters in central and peripheral vacuoles of young oogonia. Small vesicles were present closely adjacent to both the central vacuole membrane (Fig. 13) and the oogonial wall (Fig. 14). After oosphere-initial development (Fig. 16) the space between the central granule and the limiting membrane of dense-body vesicles was completely filled by electron-dense material which often showed myelin-like banding (Fig. 17) similar to that in dense-body vesicles of Saprolegnia furcata (Gay et al., 1971). The central vacuoles of oogonia appeared to be fully enlarged (Fig. 12) before the appearance of oosphere initials (Fig. 16).

The volume density of dense-body vesicles relative to the peripheral protoplasm progressively decreased from young through to oosphere initial stages, while that of lipid progressively increased (Table 1). The volume densities of mitochondria and nuclei relative to the peripheral protoplasm did not change significantly between young and late pre-oosphere initial stages despite the reduction in protoplasmic volume, but decreased between late pre-oosphere initial and oosphere initial stages (Table 1). The volume density of the matrix relative to the peripheral protoplasm showed a slight decrease between young and late pre-oosphere initial stages but this was of low statistical significance (Table 1). No significant change in matrix volume density occurred between late pre-oosphere initial and oosphere initial stages (Table 1). The volume density of central vacuoles relative to the total oogonium volume increased by 48.2% of the oogonium volume between young and late pre-oosphere initial stages (Table 1). The volume density of lipid relative to the total volume did not differ significantly between young and late pre-oosphere initial stage oogonia (Table 1). Volume densities of other organelles, and of the matrix, relative to the total oogonium volume were lower in late pre-oosphere initial stage oogonia than in young oogonia (Table 1).

The mean diameter of dense-body vesicles progressively decreased during oogonium maturation but the differences between stages were of relatively low significance (Table 2). The numerical density of dense-body vesicles relative to the peripheral protoplasm did not change significantly from young through to oosphere initial stages but decreased between young and late pre-oosphere initial stages relative to the total oogonium volume (Table 2). Of vesicle profiles that contained dense-body granule profiles, 5.4 ± 1.8% (1) contained two or more granules in young oogonia, 20.2 ± 5.3% (2) in late pre-oosphere initial stage oogonia and 16.5 ± 8.5% (3) in oosphere initial stage oogonia. Values 1 and 2 differ significantly at \( P < 0.001 \); values 2 and 3 do not differ significantly at \( P = 0.05 \).

Precise calculation of organelle volume densities relative to the total oogonium volume for oosphere initial stage oogonia was not possible since the oogonium volume fraction
Oogonia from cultures of *Saprolegnia diclina* grown continuously with 6.85 μM-CaCl₂. Bar markers represent 10 μm in Fig. 1, 1 μm in Figs 2, 3 and 5, 0.5 μm in Figs 8, 9 and 10 and 0.25 μm in Fig. 4.

Fig. 1. A median section of a young oogonium with a small central vacuole system (CV) and thick peripheral protoplasmic layer.

Fig. 2. Part of the central vacuole system (CV) and adjacent peripheral protoplasm of a young oogonium with nuclei (N), mitochondria (M), lipid bodies (L) and dense-body vesicles (DBV).

Fig. 3. A small peripheral vacuole (PV) from the peripheral protoplasm of a young oogonium.

Fig. 4. A Golgi dictyosome with associated small vesicles (SV) from a young oogonium.

Figs 5, 6, 7. Serial sections of four dense-body vesicles (a, b, c and d) from a young oogonium.

Figs 8, 9. Dense-body vesicles with two dense-body granules each from a young oogonium.

Fig. 10. Vacuoles with scattered electron-opaque granules (arrowed) from a young oogonium.
Oogonia from cultures of *Saprolegnia diclina* grown continuously with 6.85 μM-CaCl₂. Bar markers represent 10 μm in Figs 12 and 16, 2 μm in Fig. 15, 0.5 μm in Figs 13 and 14 and 0.25 μm in Figs 11 and 17.

Fig. 11. A dense-body vesicle with blebbed membrane (arrowed) and associated small vesicles (SV).

Fig. 12. A late pre-oosphere initial stage oogonium with a large central vacuole (CV) and thin peripheral protoplasmic layer.

Fig. 13. Small vesicles (SV) close to the central vacuole (CV) of a late pre-oosphere initial stage oogonium.

Fig. 14. Small vesicles (SV) close to the wall (W) of a late pre-oosphere initial stage oogonium.

Fig. 15. Part of the central vacuole of a late pre-oosphere initial stage oogonium with both dispersed and clustered electron-opaque, granular material.

Fig. 16. An oosphere initial stage oogonium with two initials.

Fig. 17. A dense-body vesicle (DBV) from an oosphere initial stage oogonium.
Table 1. Volume densities of some organelles in oogonia of *S. didinn* from high Ca\(^{2+}\) and Ca\(^{2+}\)-free culture regimes at different developmental stages

Values given are the mean and standard error of the mean and were determined from electron micrographs, unless otherwise stated. Consecutive data pairs within rows differ significantly at *P* < 0.01, unless otherwise stated. Significance levels for differences between high Ca\(^{2+}\) and Ca\(^{2+}\)-free data pairs follow the data pairs in parenthesis.

<table>
<thead>
<tr>
<th>Organelle</th>
<th>Ca(^{2+}) regime</th>
<th>Percentage volume density relative to peripheral oogonal protoplasm only</th>
<th>Percentage volume density relative to total oogonium volume</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Young, septum-</td>
<td>Late pre-oosphere</td>
<td>Oosphere</td>
</tr>
<tr>
<td></td>
<td>delimited stage</td>
<td>initial stage</td>
<td>initial stage</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dense-body vesicles</td>
<td>High Ca(^{2+})</td>
<td>31.8 ± 2.2 (0.4)</td>
<td>18.5 ± 0.6</td>
</tr>
<tr>
<td>Ca(^{2+})-free</td>
<td>29.6 ± 1.6</td>
<td>23.2 ± 0.9</td>
<td>18.7 ± 0.9</td>
</tr>
<tr>
<td>Peripheral vacuoles</td>
<td>High Ca(^{2+})</td>
<td>6.8 ± 0.6 (0.8)</td>
<td>6.8 ± 0.6† (0.04)</td>
</tr>
<tr>
<td>Ca(^{2+})-free</td>
<td>6.5 ± 0.9 (0.001)</td>
<td>5.2 ± 0.5</td>
<td>7.0 ± 1.0</td>
</tr>
<tr>
<td>Lipid bodies</td>
<td>High Ca(^{2+})</td>
<td>13.8 ± 0.5 (0.02)</td>
<td>28.9 ± 0.5</td>
</tr>
<tr>
<td>Ca(^{2+})-free</td>
<td>10.8 ± 1.1 (0.02)</td>
<td>26.0 ± 0.2</td>
<td>31.5 ± 0.8</td>
</tr>
<tr>
<td>Mitochondria</td>
<td>High Ca(^{2+})</td>
<td>3.3 ± 0.1 (0.02)</td>
<td>3.3 ± 0.1†</td>
</tr>
<tr>
<td>Ca(^{2+})-free</td>
<td>4.5 ± 0.3 (0.04)</td>
<td>3.6 ± 0.2</td>
<td>2.6 ± 0.3</td>
</tr>
<tr>
<td>Nuclei</td>
<td>High Ca(^{2+})</td>
<td>2.2 ± 0.0† (0.01)</td>
<td>2.2 ± 0.0†</td>
</tr>
<tr>
<td>Ca(^{2+})-free</td>
<td>3.3 ± 0.3 (0.03)</td>
<td>2.9 ± 0.2</td>
<td>1.0 ± 0.4</td>
</tr>
<tr>
<td>Central vacuoles</td>
<td>High Ca(^{2+})</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Ca(^{2+})-free</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Matrix</td>
<td>High Ca(^{2+})</td>
<td>42.6 ± 1.5§ (0.02)</td>
<td>39.2 ± 1.0† (0.003)</td>
</tr>
<tr>
<td>Ca(^{2+})-free</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
</tbody>
</table>

ND, Not determined; NA, not applicable.

* Means only; calculated from values relative to peripheral protoplasm using the same conversion factor as for the late pre-oosphere initial stage.
† Consecutive data pairs within rows not differing significantly at *P* = 0.4 have been combined.
‡ From light microscope measurements.
§ Difference significant at 0.05 < *P* < 0.1.
Table 2. Diameter and numerical density of dense-body vesicles in oogonia of S. diclina from high Ca²⁺ and Ca²⁺-free culture regimes at different developmental stages

Values given are the mean and standard error of the mean.

<table>
<thead>
<tr>
<th>Developmental stage</th>
<th>Mean diameter (µm)</th>
<th>Numerical density (µm⁻³)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>High Ca²⁺ regime</td>
<td>Ca²⁺-free regime</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Young, septum-delimited</td>
<td>0.861 ± 0.065*†</td>
<td>0.728 ± 0.050*</td>
</tr>
<tr>
<td>Late pre-oosphere initial</td>
<td>0.757 ± 0.011</td>
<td></td>
</tr>
<tr>
<td>Oosphere initial</td>
<td>0.697 ± 0.029‡</td>
<td>0.553 ± 0.036‡</td>
</tr>
</tbody>
</table>

ND, Not determined.

* Difference not significant at $P = 0.1$.
† Difference significant at $P = 0.02$.
‡ Data pairs within rows differ significantly at $P < 0.001$.
§ Data pairs within columns differ significantly at $P < 0.001$.
|| Difference significant at $P = 0.05$.
†† Difference significant at $P = 0.01$. 

Data pairs within columns differ significantly at $P < 0.001$. 
Data pairs within rows differ significantly at $P < 0.001$. 

Calcium and oogonia of Saprolegnia
Table 3. Effect of CaCl$_2$ concentration on oogonia and oospores of S. diclina

Cultures were grown for 5 d in medium 1 then for 10 d in medium 2 before sampling, with sampling restricted to mycelial growth made in medium 2. Values given are the mean and standard deviation for eight replicates with 125 oogonia and 40 oospheres (oospores) examined per replicate.

<table>
<thead>
<tr>
<th>CaCl$_2$ concn (μM)</th>
<th>Medium 1</th>
<th>Medium 2</th>
<th>Aborted oogonia (%)</th>
<th>Aborted oospheres (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>6·85</td>
<td>6·85</td>
<td>0·7±1·1*</td>
<td>47·7±12·3†</td>
<td></td>
</tr>
<tr>
<td>1·71</td>
<td>6·85</td>
<td>2·1±1·7*‡</td>
<td>50·6±8·0‡‡</td>
<td></td>
</tr>
<tr>
<td>1·71</td>
<td>0</td>
<td>52·9±14·8‡‡</td>
<td>97·2±5·0‡‡</td>
<td></td>
</tr>
</tbody>
</table>

* Difference not significant at $P = 0·05$.
† Difference not significant at $P = 0·5$.
‡‡ Difference within columns significant at $P < 0·0001$.

occupied by the irregularly-shaped, central vacuole could not easily be measured. Table 1 gives mean values for volume densities in oosphere initial stage oogonia relative to the total oogonium volume determined using a conversion factor derived from measurements on late pre-oosphere initial stage oogonia. If a further increase in the volume fraction occupied by the central vacuole occurs between late pre-oosphere initial and oosphere initial stages, these values will be overestimates.

Ca$^{2+}$-free regime

Almost all oogonia formed in cultures grown under the Ca$^{2+}$-free regime eventually showed abnormalities detectable by light microscopy. The percentage of oogonia that aborted (oogonia forming no recognizable oospheres or oospores) was increased about 50-fold relative to that in cultures grown continuously with high (6·85 μM) Ca$^{2+}$ (Table 3). In the remaining oogonia that did form oospheres, the percentage of oospheres that aborted (oospheres or oospores not showing normal maturation; Fletcher, 1978) reached almost 100% (Table 3). Initial growth in low Ca$^{2+}$ medium did not significantly increase abortion rates when followed by transfer to high Ca$^{2+}$ medium (Table 3).

Aborted oogonia showed extreme protoplast degeneration (Fig. 18). Crystalline inclusions were present in central vacuoles of unaborted oogonia (Fig. 20) but organelles of unaborted oogonia appeared normal (Fig. 19). About 50% of the oogonia examined at late pre-oosphere initial stage had failed to form secondary wall layers (Figs 21, 22) and had small, cytoplasmic vesicles close to their walls (Fig. 22).

Volume densities of dense-body vesicles and of peripheral vacuoles were unaffected by growth under the Ca$^{2+}$-free regime in young oogonia but the volume density of dense-body vesicles was increased in late pre-oosphere initial stage oogonia and the volume densities of both organelles were increased in oosphere initial stage oogonia (Table 1). Volume densities of mitochondria and nuclei were increased in young oogonia but were unaffected in late pre-oosphere initial and in oosphere initial stage oogonia (Table 1). The density of lipid was reduced at all three developmental stages (Table 1). The mean diameter of dense-body vesicles was reduced in late pre-oosphere initial and in oosphere initial stage oogonia but was not significantly affected in young oogonia (Table 2). The numerical density of dense-body vesicles increased progressively during oogonial development and was greater at all stages than in oogonia from cultures grown under the high Ca$^{2+}$ regime (Table 2). The percentage of vesicle profiles that contained dense-body granule profiles having two or more granule profiles each in young oogonia (5·1±2·3) did not differ significantly ($P > 0·5$) from that in young oogonia grown under the high Ca$^{2+}$ regime. The percentage in oosphere initial stage oogonia (8·5±3·4) was significantly lower ($P < 0·01$) than in high Ca$^{2+}$ regime oogonia at the same developmental stage.
Calcium and oogonia of Saprolegnia

Oogonia from cultures of *Saprolegnia diclina* either grown for 5 d with 1.71 µM-CaCl₂ then in CaCl₂-free medium for 5 to 10 d (Figs 18, 19, 20, 22) or grown continuously with 6.85 µM-CaCl₂ (Fig. 21). Bar markers represent 2 µm in Fig. 18 and 1 µm in the remaining figures.

Fig. 18. Part of an aborted oogonium.

Fig. 19. Protoplast detail from an unaborted, late pre-oosphere initial stage oogonium with nucleus (N), mitochondria (M), dense-body vesicles (DBV) and lipid bodies (L).

Fig. 20. Crystalline inclusions in the central vacuole (CV) of an oogonium (section unstained).

Fig. 21. Primary (P) and normally developed, secondary (S) wall layers of a late pre-oosphere initial stage oogonium.

Fig. 22. Peripheral zone of a late pre-oosphere initial stage oogonium with primary wall layer only and small vesicles (arrowed) close to the wall.

**DISCUSSION**

The presence of similar clusters of granular material in central and peripheral vacuoles of young, septum-delimited oogonia and in the central vacuoles of late pre-oosphere initial stage oogonia suggests that the large central vacuoles of late pre-oosphere initial stage oogonia form by enlargement of the small central vacuoles of young oogonia and that some peripheral vacuoles of young oogonia may become incorporated into the central vacuoles. The decrease in volume density of dense-body vesicles during oogonial development suggests transfer of dense-body vesicle contents into the developing central vacuoles.
Direct fusion of dense-body vesicles with the enlarging central vacuoles has been suggested for central vacuole development in oogonia of *S. ferax* (Gay *et al.*, 1971), but no evidence for this, such as fusion profiles or incorporation of dense-body granules into central vacuoles, was observed in *S. diclina*. The increase between young and late pre-oosphere initial stages in the percentage of dense-body vesicles having two or more dense-body granules suggests that dense-body vesicles in *S. diclina* may fuse mutually during oogonium development. Mutual fusion of dense-body vesicles has been reported to occur at a later developmental stage, during maturation of cleaved oospheres into oospores, in *S. terrestris* (Howard & Moore, 1970) and *S. furcata* (Beakes & Gay, 1978). Mutual fusion of dense-body vesicles in oogonia of *S. diclina* could at least partially account for the reduction in numerical density of dense-body vesicles relative to the total oogonium volume between young and late pre-oosphere initial stages. Such fusion should also increase the dense-body vesicle size, but no such increase was observed. Although some of the small, cytoplasmic vesicles seen in oogonia of *S. diclina* are probably Golgi-derived vesicles that contribute to oogonial wall development, as in oogonia of *S. furcata* (Heath *et al.*, 1971; Gay *et al.*, 1971), the blebbed appearance of the limiting membranes of some dense-body vesicles in *S. diclina* suggests that some small vesicles may be formed by budding from dense-body vesicles, thus reducing the dense-body vesicle size. Fusion with the vacuolar membrane of small vesicles derived from dense-body vesicles could provide a mechanism for transfer of the dense-body vesicle membrane and contents to developing central vacuoles alternative to that of direct fusion of entire dense-body vesicles with central vacuoles. The presence of small vesicles close to the central vacuole membrane is consistent with this. The total decrease in mean volume densities of organelles relative to the total oogonium volume between young and late pre-oosphere initial stages (26.6%) is insufficient to account for the increase in mean central vacuole volume density (48.2%). Additional reduction of protoplasmic volume density possibly occurs by transfer of water from the cytoplasmic matrix to developing central vacuoles, thus reducing the matrix volume.

The appearance of late pre-oosphere initial stage oogonia and of oosphere initial stage oogonia suggests that, at least in *S. diclina*, oosphere initials form by the drawing into heaps of a peripheral protoplasmic layer after central vacuole enlargement. Fletcher’s (1978) time-lapse study of living oogonia of *S. diclina*, although showing a diffuse appearance to the central vacuole outline shortly before the formation of oosphere initials, supports the heaping hypothesis (DeBary, 1881; Hartog, 1895; Trow, 1899) rather than the cleavage hypothesis (Howard & Moore, 1970; Gay *et al.*, 1971) for at least the major part of the process of oosphere initial formation.

The protoplasts of oogonia of *Saprolegnia* spp. are cleaved into oospheres by eventual fusion of central vacuole membranes with the oogonium plasmalemma (Howard & Moore, 1970; Gay *et al.*, 1971). As pointed out by Gay & Greenwood (1966) for sporangia of *S. ferax*, where a similar cleavage process occurs during formation of zoospores, fusion of central vacuole membranes with the plasmalemma membrane, with the central vacuole membrane becoming incorporated into the plasmalemmas of the cleavage products, suggests a similarity in structure between the central vacuole and plasmalemma membranes. The plasmalemma membrane in fungal hyphae, including hyphae of Oomycetes, is probably derived from Golgi cisternae (Girbardt, 1969; Grove *et al.*, 1970; Heath *et al.*, 1971). Cleavage vesicle (vacuole) membranes which eventually fuse with the plasmalemma membrane may be derived from Golgi cisternae in sporangia of *Phytophthora parasitica* (Hohl & Hamamoto, 1967) and *Thamnidium elegans* (Fletcher, 1973). The enlargement of the central vacuoles (cleavage vacuoles) in oogonia of *S. diclina* by fusion with them of small vesicles derived from dense-body vesicles suggests a radically different mode of cleavage vacuole development, unless dense-body vesicles themselves are ultimately derived from Golgi dictyosomes.

Data presented here (Table 1) show that except for central vacuoles and lipid bodies,
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Organelle volume densities relative to the total oogonium volume decrease during oogonial development. The decrease in nuclear volume density is consistent with the well-documented breakdown of supernumary nuclei during oogonial development (Dick & Win-Tin, 1973). The increase in nuclear volume that occurs in the early stages of nuclear division in oogonia of Saprolegnia spp. (Howard & Moore, 1970; Beakes & Gay, 1977) is possibly completed and the breakdown of supernumary nuclei already well advanced in S. diclina by the late pre-oosphere initial stage. A reduction in the number of mitochondria, possibly associated with the onset of dormancy, occurs during oospore maturation in S. furcata (Beakes & Gay, 1978). The reduction in the volume density of mitochondria relative to peripheral protoplasm between the late pre-oosphere initial stage and the oosphere initial stage in oogonia of S. diclina is possibly a preliminary stage of a similar reduction of the number of mitochondria in this species.

Quantitative effects of Ca²⁺ deficiency on dense-body vesicles reported here – increased volume density at late pre-oosphere initial and at oosphere initial stages, increase in numerical density between young and late pre-oosphere initial stages and the reduced percentage of dense-body vesicles with more than one dense body granule in oosphere initial stage oogonia – suggests that transfer of dense-body contents to developing central vacuoles and mutual fusion of dense-body vesicles may be partially inhibited by growth under Ca²⁺-deficient conditions. It has been suggested that dense-body granules are composed of phospholipid (Gay et al., 1971; Gay, 1972) and that they may be important suppliers of endogenous energy during periods of high cellular activity such as oogonial cleavage (Beakes & Gay, 1978). If this is so, and if dense-body vesicles are involved in development of the central vacuole in oogonia of S. diclina as proposed here, any disturbance of normal dense-body vesicle behaviour, such as that observed here under Ca²⁺-deficient conditions, might be expected to seriously affect oogonial development. Wall development in oogonia of Saprolegnia spp. is thought to involve wall vesicles (Gay et al., 1971; Heath et al., 1971), similar in appearance to the small cytoplasmic vesicles found adjacent to oogonial walls of S. diclina, that release their contents by fusion with the oogonial plasmalemma. Ca²⁺ deficiency inhibits exocytotic release of secretory vesicle contents in a range of cell types (Douglas, 1974). The release of wall-vesicle contents involved in oogonial wall synthesis might be similarly inhibited by Ca²⁺ deficiency in some oogonia, inhibiting normal secondary wall development. Absence of a significant effect of Ca²⁺ deficiency on nuclear and mitochondrial volume densities in oosphere initial stage oogonia might indicate an ability of the mechanisms that control quantitative changes occurring in normally developing oogonia grown in high Ca²⁺ medium to compensate for initially aberrantly high nuclear and mitochondrial numbers in oogonia from cultures grown under conditions of Ca²⁺ deficiency. Crystalline inclusions in central vacuoles of oogonia might be present in crystalline form in living oogonia or might represent soluble-phase vacuolar components deposited as crystals during specimen preparation.

Mitochondrion-controlled cytosol Ca²⁺ concentration is believed to be involved in the control of cell activity through a wide range of Ca²⁺-dependent functions, including activation or inhibition of enzyme systems, activation of microfilaments, functioning of cyclic nucleotides and several membrane-linked functions (Carafoli, 1974; Berridge, 1976; Bygrave, 1978). Ca²⁺ deficiency can inhibit pyruvate dehydrogenase-mediated formation of acetyl-CoA required for fatty acid and lipid synthesis (Randle et al., 1974). Inhibition of pyruvate dehydrogenase activity by Ca²⁺ deficiency in S. diclina could affect synthesis of dense-body granule phospholipids and might also account for the observed reduction in volume density of lipid bodies in oogonia from cultures grown under conditions of Ca²⁺ deficiency. Ca²⁺ deficiency might inhibit uptake of amino acid constituents of the growth medium by disturbing the normal cell membrane integrity and function in S. diclina, and hence inhibit protein synthesis, as suggested by Cameron & LeJohn (1972) for Achlya. Probably, the overall effect of Ca²⁺ deficiency on oogonia of S. diclina is the cumulative...
result of the disturbance by Ca\(^{2+}\) deficiency of a wide range of Ca\(^{2+}\)-dependent functions, many not yet investigated in *Saprolegnia* or related genera.

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